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M. P. WOO

Member of the American Society of Mechanical E for the Advancement of Science; tone Empireer, Supersulendent of Mo

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PREFACE.

Since the publication of the papers from and Steel," in the years 1894-1901, the many requests from engineers and others there available form than in the Transa Society of Mechanical Engineers and in the American and foreign technical journals. I mainly rewritten and new matter added to The characteristics of oils, pigments, and pacipal protective contings for terric and other at length under their respective chapters.

author that the collected data will afford formation of what paints are composed and of them. The technical journals have give to the subject of proctective coatings for a not always available when comparison with



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Paint Statistics and General Character.

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Iron, Iron-Oxide, Copperas, Ochre-



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RUSTLESS COATINGS.

CHAPTER I.

PAINTS: OF WHAT COMPOSED, HOW DESTROYED, CLASSIFICATION AS TRUE PROMENTS AND INERT SUBSTANCES, ADULTERANTS, ETC.*

What is paint? This question can be answered in a broad way by saying: It is any liquid or semi-liquid substance applied to any metallic, wooden, or other surface, to protect it from corrosion or decay, or to give color or gloss, or all of these qualities, to it.

A better definition would be, that paint is a compound of a pigment and a liquid, usually applied to any surface with a brush, for the purpose of protection, or to secure artistic effects; which liquid, after undergoing certain changes, in part mechanical, or chemical, or both, has the power of holding the pigment to the coated surface. It is evident that the latter definition would also include those compounds which are applied to many surfaces either hot or cold as a bath, rather than by a brush, solely as a matter of convenience or rapidity; and particularly so when metallic members of large size, or with intricate and hidden parts, are to be protected.

The essentials of a good paint, for whatever use intended, are:

First. That it shall achieve firmly to the surface over which it is spread, and not chip or peel off. It must be non-corrosive to the material it is used to protect, as well as to itself under long periods of atmospheric exposure and chemical changes. It must form a surface hard enough to resist frictional influences, yet elastic enough to conform to all changes of temperature, or with a coefficient of

influences that govern resums.

Third.—That it shall dry with sufficient rapidity. This function depends mostly upon the vehicle or liquid used with the pigment, though the pigment has in many cases an influence, as will be seen further on.

Fourth.—That it shall have proper durability, which is a function both of the pigment and liquid. And as the question of cost is in many cases the governing factor in the selection of a paint, the question of durability may be regarded as the most important one of the list. It should be understood, however, that a paint can be durable per se, and not be protective in the strict sense of the word, as can be illustrated in the case of a good paint applied to the surface of a sheet of iron conted with rust. The liquid element in the paint will not absorb or neutralize the corresion which it covers, but will dry regardless of it, and permit the destruction of the metal to progress beneath its coat.

Fifth.—Covering power, by which is meant the power of a pigment so to cover the surface to which it may be applied that its presentection from decay is not only assured, but that the minimum amount of paint shall effect this purpose,

The covering power is also used to express the power of a pigment to protect the oil from decay, in which case a large amount of pigment and a small amount of oil are used. This description of point drien more or less "flat," the pigment being exposed to the weather and held in place by the thin film of oil. It is thought by many master painters that this is the most durable and best paint for general use. On the contrary, paints that dry with a gloss have a large amount of oil and a small amount of pigment, in which case the oil covers and protects the pigment.

It may be used to express the amount of color upon the surface; as, generally, if a surface has plenty of color upon it the covering power is said to be good. To illustrate this definition: If an iron-oxide paint is proportioned so that the ratio between the pigment and the oil is by weight 50 per cent of pigment and 50 per cent of oil

when the paint is ready for spreading, and the pigment consists of 30 to 40 per cent of iron oxide, the covering power will be said to be good; but if the same proportions of 50 per cent ratio between the pigment and the oil be had, in which the iron oxide is only 5 per cent of the pigment, the covering power would be called poor; and so it would be in the case where 10 per cent of pigment and 90 per cent of oil were used. If in the two latter cases the oil contained large or liberal amounts of volatile diluents, the appearance of the surface would indicate a deficiency in the covering power of the paint.

The covering power is also commonly expressed in the amount of surface which a given weight of paint will cover. A good iron-oxide paint will cover nearly twice as much surface as white or red lead. The specify gravity of the paint also is to be considered in the definition of this power. The lightest paints have the most covering power. White lead is about 6.4 times as heavy as water; iron oxide 5 times; yellow other 3 to 4 times, etc., etc. With this variation it is manifestly almost an impossibility to get the same number of particles of the same size out of the same weight of different materials.

Fig. 1 represents the covering power of a number of paints and inert pigments.

The refracting power of light has much to do with an understanding of this covering power of paint. The greater the refracting power of the pigment is over that of the oil, the better will be the covering power. The index of refraction of air is 1 degree; water, 1.34; linseed oil, 1.48; glass, 1.50 to 1.55; silien, 1.55; feldspar, 1.54; whiting, 1.65; chrome-yellow, 3.00; vermilion, 3.20, etc. There is no exception to the rule that the finer the state of division to which any pigment is reduced, the better will be its covering power. Sulphate of lime, barytes, feldspar, silien, tale, whiting, etc., are all of low refractive power, and of themselves, independent of this refractive quality, do not constitute good pigments, though when mixed with the metallic pigments and ground together in the oil the result is a pigment of good covering power, almost as good as the best one of the combination. For instance, 80 per cent of subhate of lime

the use of which is objectionable so far as durability and protective

power are concerned.

The covering power is due to two qualities. For instance lime whitewash has very little covering power until it becomes dry Barytes covers well as a water paint, because the water leaves it as a dry powder on the coated surface. But barytes covers paorly in

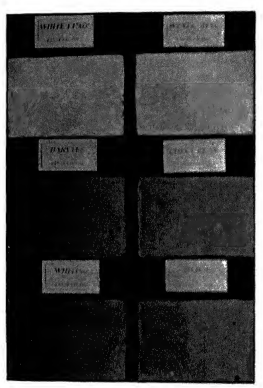


Fig. 1.-Covering power of paints.

oil, because the oil remains with it, and the light remches it through the transparent film of oil.

Prof. Von Bezold's experiments,* from which I quote, illustrate how lime, barytes, white lead, and other crystalline pignishts, when

the powder will appear white, and it will be impossible to see through it, but as soon as water is poured into the tube, the powder, to a certain degree, becomes translucent. By substituting turpentine for the water, the degree of translucency is materially increased. If a small quantity of sulphuret of carbon is added to the turpentine, a liquid is obtained that reflects (bends) the light about as powerfully as glass. If some of this liquid be poured upon the powder in the test-tube, the powder will disappear, and light passes through the tube as freely as though no powder were present.

If a solid glass rod be immersed in such a liquor (or a mixture of olive-eil and oil of cassia), it will appear as if the rod only reached to

the surface of the liquid. Within the liquid the rod cannot be seen (Fig. 2).

It is shown by these experiments that the presence of one transparent body within another is only detected by the eye when the two differ in their power of refracting light.

Many white substances are white

A white lily is white because it consists of little cells which reflect all kinds of light, again and again, until it reaches the eye from some part of its surface. Water becomes white when it is broken into

because they are in fine particles.



Fm. 2.

fine drops, as in a waterfall or on the crest of waves. White lead and zine owe their whiteness to their dense, fine, powder-like condition, and transparent glass becomes white when finely pulverized."

As stated before, the finer the pigment is subdivided, whether as a paste which is afterward thinned with oil or volatiles to a consistency to spread with a brush, or is ground in the oil direct (a process that all pigments will not endure without injury to their color—the scarlet lead chromate, for instance) to the proper consistency to

sulphate of line by weight, give most excellent in the more experiment of line by weight, give most excellent in the more experiment of the first of the be called "the fugitive colors," on account of them to account to the out, possess the light-dispersing power which decrease the covering power, when in reality for a trade excellent the first substances they are absolutely worthless. The modest form in the more more more excellent than paints; for generally the set may an

substances they are absolutely worthless. The reads of small leads nominated stains rather than paints; for generally the self-making of protection from decay or corrosion which accompany from the is solely from the oil or liquid with which the redor a first of the designing of a paint, for whatever property leads out in cludes the qualities already mentioned, vizit soils now as it into

The designing of a paint, for whatever proposed have such an eludes the qualities already mentioned, vizit, also not will the ticity, working qualities, drying qualities, durability, and will be vide one sidered later, as well as what pigments to use for the interaction of All pigments do not contain all of the above qualities. The specific naturally arises: Is it necessary for a pigment to be pure at time to with inert substances, or can a certain amount of the above by terms.

with the pigment without detriment to it?

Experiments of long duration lead to the conclusion of the oxides of iron, lead, manganese, and other strong process, and other strong process, as the mixed with large amounts of these inert substance with a form of and generally to the manifest improvement of the part of action tective agent on many structures, notably moder, to action tective agent on many structures, notably moder, to action of iron is one of the strongest of pigment in received process. It one ounce of this pigment be spread in two contractions are related as the polynomial the job be declared a satisfactory one so far accordingly action to the declared manifest of the contraction of the same real action to mixed with three ounces of barytes, knoling graphical actions as a mixed with three ounces of barytes, knoling graphical actions as a mixed with three ounces of barytes, knoling graphical actions as a mixed with three ounces of barytes, knoling graphical actions and action of the same real action to the same real actions and the same real actions to the sam

cases; but in one case there is four times as much pigned to an elementer, and in the second case three-fourths of the point see I character material. For railway cars and wooden attractions to be billity of these paints would be in favor of the second case of the second case.

paint be spread over two square feet of surface at before 16.2 and 10.0 that the amount of color per unit of surface will be 13.2 and 12.2 and 13.2

It induces and promotes oxidation in all organic substances with which it is brought into contact, and in nearly all metallic bodies. In an oxide-of-iron paint it is the oil which decomposes, it being the organic matter. The decomposition is due to the exposure to the elements aided by the oxidizing power of the oxide of iron pigment mixed with the oil. This statement holds true only where there has been no chemical change or combination between the pigment and the liquid.

Whiting, sulphate of lime, barytes, kaolin, silica, feldspar, and tale are the principal inert substances used in pigments. Whiting, gypsum, and barytes are the best of the list; the others, grinding greasy, are hard to grind, or of a nature readily decomposed by water, are objectionable. Barytes, from its great weight, is objectionable as a paste or prepared paint. Its use as an adulterant is given in Chapters VI and XVIII. The sulphate of lime (gypsum) is no doubt the best of the inert substances to mix with any pigment, all things considered. It should be thoroughly hydrated. As high as 45 per cent by weight of this substance can be mixed with 50 per cent of sesquioxide of iron for a pigment. Many of the oxide-of-iron paints are made by ignition of copperas, and a notable amount of sulphuric acid is usually left in the oxide which the heat has failed to drive off. From 2 to 5 per cent of carbonate of lime is added to neutralize the free acid, changing it to sulphate of lime. In these proportions, the pigment really consists of 50 per cent of oxide of iron and 50 per cent of inert material, all by weight. Any oxide-of-iron paint which contains hydrated oxide or free SO, will deteriorate rapidly by oxidiz-

ing the liquids, while any free SO₂ will retard the drying of the paint. A good paint prepared for spreading in ordinary temperatures upon wooden or composite structures has the ratio by volume of about one-third pigment and two-thirds oil or liquid. The practice upon one of the leading railways of the United States, where the materials purchased for paints amount to over \$300,000 yearly, is to allow 75 per cent of pigment and 25 per cent of oil by weight, for the paints applied to cars and wooden structures.

Experiments determine that the most durable paints are those which contain a large amount of pigment per unit of surface; and that

ical injury; second, the action of deleterious games, think, chemical action between the pigment and the vehicle or liquid, fourth, chemical action between the body covered and the paint, either the pigment or the liquid; fifth, the action of light; sixth, proling, recently, destruction by cleaning; eighth, water.

Many master painters and manufacturers claim that the destruction caused by cleaning and the action of water are the world of the
above causes. This is true so far as paint applied to mostler denotines
is concerned, but has no relation to the causes which effect the destruction of paint applied to iron or steel structures. As most of the
above destructive agents are common to all structures passible,
metallic, or composite) which depend in a greater of less degree for
their preservation from decay or corrosion, upon paint (under which
name all paints, oils, varnishes, japans, and surfacets are classify it
may not be amiss to discuss briefly each of these causes in detail before
citing the destructive agencies which relate solely to the corrosion
of metallic structures, the prevention of which will require the consideration of other preservative methods than paints, or which may
be used in connection with paint to secure the best protective teerible.

First.—Mechanical injury to wooden structures is not a section

First.—Mechanical injury to wooden structures is not a settlere cause of deterioration of paint. Near the sensions the wind and said have the effect of a sand-blast, which cuts away the paint rapedly, and in this case the more elastic the paint, the less will be the mechanical injury. This sand-blast action is quite as effective on form structures, and as generally they are of a more important character than the wooden cottages and minor buildings on the paint character than the wooden cottages and minor buildings on the paint of a soft, spongy nature it will resist the sand-blast, but will absorb metaltice from the air, and hasten either the oxidation of the paint of the metallic surface which it covers

A further injury to metallic structures can be characterized the head of mechanical, viz.: that arising from the expansion and contraction of the various parts from the atmospheric charges which are constantly going on, changes ranging from 40 degrees 1 to 150 degrees F. not being unusual. It is an impossibility to proportion a paint compound so that its coefficient of clasticity will be the contract.

is the tale of the salesman. It may be argued that, these changes coming from the external surface of the paint and being transmitted through its coating, it will be the first to adjust itself to the new or varying relation between the metal and the paint, and so will work to the advantage of the paint in making the change, this being in ordinary cases a gradual one. If the paint is of an elastic, close-elinging material, and not a hard, vitreous one, the claim will hold good.

The compounds which most closely partake of this mature, will

be spoken of hereafter. An addition to this problem will be had when the strains due to the action of wind, the passage of railway trains, and those due to changes of a sudden and vibratory character, together with the action of snow, hail, and water driven at high velocities, are added to the temperature changes. These strains necessarily come to the metal first, and whatever changes occur in the bars by the strain, the paint must accompany them. As these strains are generally of a vibratory or percussive character, it can easily be seen why they should be classed in the list of mechanical injuries. In fact, they are a succession of blows which the structure must absorb, withstand, and extinguish within itself or its connections; the structure then returning to its normal condition. the paint or other protective covering must accompany it, instead of loitering by the way and being grounded or "left" in the chain of operations. Second, The action of deleterious gases is very familiar to those

Second,—The action of deleterious gases is very familiar to those who have studied paints and protective compounds. Sulphuretted hydrogen is one of the most common and active of these gases, and is formed in excessive amounts wherever coal is distilled for illuminating gas. Sulphurous acid fumes also, being disengaged in the combustion of coal in the many arts, transportation, and manufacturing processes of the day; gases engendered in workshops, being of a compound character carrying ammonia, carbonic acid, nitric acid, and other fumes, are active agents of corrosion to metallic bodies, also to the paint compounds that cover them. (See Analysis of Smoke, Chapter XXXVI.)

Third,—Chemical action between the pigment and the vehicle. This is an exceedingly important field of inquiry, and largely an

ricent. Terring. Taunts vary greatry in each power to admere to either metallie, wooden, or other surfaces; notably zinc white, which peels under almost any condition or from any surface to which it umy be applied. There is no other pigment which possesses this property in so marked a degree, and it is difficult to assign any reason why it should peel so budly. A possible cause is that the zine white combines with the oil used in the paint and forms one of the compounds known as metallic soap, this particular one being zinc soap, a hard, brittle, non-adhesive substance, easily removed by mechanical injury, water, and in the process of cleaning, etc. Galvanized iron possesses the property of causing almost any paint applied to its surface to peel; in fact, it is one of the worst substances to cover with a pigment in a satisfactory manner. Experiments made by a leading railway company in the United States, in which a number of the best pigments in use by that company for all descriptions of railway work were tried upon galvanized-iron car-roofs and other galvanized work, cornices, etc., showed at the end of three years that but one of the list was in any manner satisfactory, and this one was a patented compound with bisalphide of carbon as the vehicle, Ordinary trade colors are of the most unreliable nature when applied to galvanized iron expose I to the trying conditions of railway service. Various reasons have been given for this peculiar action of paint upon galvanized iron. One of the most plausible is that the use of sul-ammoniac in the process of galvanizing causes the formation of a thin film of the basic chloride of zinc on the surface of the metal being galvanized, which material, being of a hygroscopic nature. acts as a repellant to prevent the close adherence of the paint to the metal, and the pigment dries as a skin over it. Sheet zinc does not hold some kinds of paint. Sheet lead also is difficult to cover, and paints which take tin and lead will not always adhere to zine. As a

general rule, the strong oxide paints take these metals better than tale, other, and the earthy pigments. No positive general statement can be given, and the problem of the adaptability of paint to a metal to prevent pecling still needs study for each application. Paints applied in cold weather, and which are exposed to a frost while drying, will always peel, unless the paint is warmed to about 120 degrees F.

elements of the liquid, beginning to dry from the outside surface, forms a thin but hard or vitreous surface which retards the further evaporation of the volatiles and prevents the access of oxygen from the air, which is necessary in the process of drying. If the surface thus covered has been painted while at a low temperature or during a damp or foggy atmospheric condition, and ason after there is a marked rise in the temperature or a fall in the hygroscopic condition of the atmosphere, then the paint is liable to peel at once, or ason after the change. This effect is hastened where the coating is a heavy one, or one hard to spread by reason of the earthy or inert substances in the pigment, or if benzine has been used as a drief.

As a general rule, the more substances that enter into a point, either as pure pigments, inert substances, or in the composition of the liquid, the more liable it is to peel. A small amount of tab or animal or non-drying vegetable oils, though exidized by the addition of metallic salts and used in connection with lurged or other sicentise oils, also hastens and provides for the certainty of the peeling.

The preling of paint from wooden surfaces is very common, particularly if applied on unseasoned lumber that contains most are

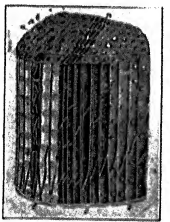


Fig. 3. -Cellular formation of wood causing the peeling of

and air in the cellular formation of the wood as shown by the cut. The air and moisture in the cells expand upon a slight rise in temperature, and in their efforts to escape through the dreed pointskin, push it up in the form of blisters that contain the condensed moisture, and results in the preling of the paint in blisters or in strips.

A pigment composed of a number of substances, the different materials of which by themselves would form the basis of a good paint, when combined together with the liquid, necessarily undergo a different chemical action than the several members of the pigment.

first to break down or change, the decay of which hastens the decomposition of the others and releases the bond between the paint and the surface over which it is spread, and the peeling process is effected.

That these chemical changes exist in the above stated case cannot be denied, but have not been well accounted for. The fact remains, however, that certain paints peel, and though analysis of the peeled portion may reveal nothing to indicate the reason for the peeling, it is seldom possible to get a sample of the original paint applied, to compare its constituents with the peeled sample, and the cause is relegated to the hidden drawer of the paint-shop, near which some scapegoat can be found to bear the burden of failure. (For other notes on the peeling of paint, see Index.)

Seventh.—Destruction by cleaning. This cause of the deteriora-

tion and destruction of paint applies more particularly to wooden structures, railway cars, and kindred objects, than to those of a metallic character. It may be sufficient to say we do not wash down an iron bridge, roof-truss, or steamship, with a view to its presenting a clean face for inspection and painting. Almost all the binding materials of dried paints and varnishes are more or less acted upon by caustic and carbonated alkalies, and but little of the soap in the market is free from these substances. The detergents sold for cleaning are all mixtures of sal-soda and caustic substances with lime, pumice, and other inert materials, and the more effective they are for removing dirt, the better they are for the destruction of the paint. If, in the economy of domestic household matters, two removals are equal to one fire, then it may be cited with equal force that two good scrubbings with any washing compound, and most of the soaps of commerce, applied with a stiff brush, will be equal to the next painter's bill to restore matters to their pristine state. Aside from the element of cost, it is no doubt the better practice, so far as the ultimate preservation of any metallic structure is concerned, that it should be washed clean with some of the detergent compounds of the day, in a very weak solution to remove the dirt, then sponged with a liberal amount of clean water, then be allowed to dry thoroughly before the new paint is applied; but I must confess as an engineer, that the above method of painting is rare, and that the rule is for the paint to be put on regardless of cleaning the old coat, and

Eaglith, Water, The destructive action of water upon paint applied to structures of any material, either upon their internal or external surfaces, is very strong, and will rank next in destructive qualities to the detergent soap and scrubbing brash. Inside painting lasts longer than outside, principally because it is less exposed to the action of water. Direct experiments show that sixed insert and other sicentive rule, without pagment, are not residual or unter-repellent. When the oil is well street, the application of water always causes the cal to assume a shrivefled appearance, showing that it has absorbed mosture and expanded, and dismingration has commenced. If the exposure by long continued, the whole conting of dried oil will slump news from the surface over which it to opposit Rnin-water, from the sensible amount of amounts that it carries, improposed this destructive action on the drivit oil, and the show wasting away of good paints containing juginents less known to resist aging influences, and which have been hardened by time, can be attributed to this action.

The ordinary test by master painters, of the ability of an oil or paint to resist moisture is to cont a surface, usually of glass, and when well dried, to immerse it in water for a few hours and note the changes in color and integrity of the paint

Or. Dudley's experiments for the Pennsylvania Hadroad on the action of water upon paints, are interesting from the rate which was exercised in making them and recording the results. Several camples of a paint designed for use upon cars and weeden structures were unde with raw lineard-oil and a very small amount of japan, the same liquid being used for all the samples with varying amounts of pigment, all the proportions being by weight. Two coats of these paints were spread upon glass, and allowed to harden for two to three weeks. These samples were then placed side by side, and a small portion of the surface of each covered with a globale of nater. This globale was covered to prevent evaporation, and then allowed to stand for twelve to fourteen hours.

No. 1 was the linear sile and income at any

linseed-oil and the pigment has the specific gravity of ordinary oxideof-iron paints.

At the end of the period named, the behavior of the samples was as follows: No. 1 coating was found to have cleaved off the glass and had become shrivelled wherever the water had touched it. Apparently the dried linseed-oil had soaked up water, much as a sponge acts as an absorbent. On allowing the water to evaporate, the coating dried down again, but not uniformly, and was apparently weakened in texture.

No. 2 showed the same characte istics.

No. 3 showed the same, but in a less degree.

No. 4 did not cleave off the glass, but showed where the water had stood.

No. 5 showed a spot in the same way, but in a less degree than No. 4.

Nos. 6 and 7 showed but very little action.

It can be noted that here linseed-oil dried for some two months absorbed less water than freshly dried oil, while very old dried oil lost this absorbent quality and became almost water-repellent.

To successfully design a paint which will resist all of the previously named destructive agencies, is a difficult matter. The field is an enormous one to cover and but little positive knowledge has yet been obtained, though the investigators and experiments have been legion, and the literature on the subject embraces volumes. Time is an essential factor in the test of the qualities of a paint, and if the experimenter is required to wait five or ten years to determine the merits of any paint, or what effect a slight modification of the proportions has upon any one or more of the eight destructive agencies heretofore stated, a life could be spent and possibly no conclusion reached.

Experiments are numerous in the field of designing a water-proof coating to be applied over the pigment which has been found to possess the most preservative qualities, independent of the water-repellent features, but the goal is not yet reached. How effectually a thin coating of the proper material can protect the surface of a paint which it covers, can be seen in the lettering of old sign-boards, which is perhaps an example of the most durable paint of which we have any record

lampblack is one of the best water-repellents known, that it is practically indestructible by exidation or acids, and being per se of an only or greasy nature, when mixed with a pure of theseed in these cases), and being in a measure clastic, it has effectually preserved the surfaces and not allowed the water to reach the underlying coats of white lead.

Having set forth the general character of what a paint should be for the purpose of protecting structures from decay or corresion, and having indicated the most effective causes which provoke or promote the destruction of the object and its protector, it may not be amiss to speak more definitely upon those materials which enter into paint compounds which vield the best results in general practice. These results are based upon the experience thus far at hand as recorded or accepted data, and not the hypothesis of some persons or persons whose single or joint lives may be two short a period, as compared with the life of the structure they are striving to protect from decay, to realize the meritorious features of their experiment.

CHAPTER II

STATISTICS AND GENERAL CHARACTER. PAINTS:

THERE are in the United States at the present date (1903) about 420 firms engaged as manufacturers and compounders of pigments, pastes, and paints of all grades, representing a yearly output equivalent to about 90,000,000 gallons of mixed paints, that cost not far from \$65,000,000.

This represents about 570,000 short tons, and would cover with one coat 900,000 acres or 1400 square miles of surface, requiring 50,000 painters to spread it.

The following details are the average amounts of the principal pigments used in the United States for the years 1898 to 1902;

	•						
Tron oxide, 23,500 short tons. Value \$10.7.	5 to \$1 U for	1.00	Į×:1	ton			
AND IN A COLUMN THE RESERVE A COLUMN TO A COLUMN TO A COLUMN THE RESERVE A COLUMN TO A COLUMN THE RESERVE A COLUMN	***			fi if 24,			
White lend ground in oil, 85,100 short tons.	Value	-5.25	10	5.50	cents	ter	nannd.
" " dry25,100 " "	**	4.70	111	a on	**	1 11	- 11
" " imported 300 to 700 tons,		***	***	9,2461			
Red lend 11,100 short tons,	11	5.30		8.80	11	**	
" " imported 400 to 800 tons,		11.1917		13,171		•	
Litherge	11	5.30	+	a un	11	**	**
" imported Itt to 350 tons,		******		13,/76.1			••
Orange mineral 10,200 tons.	11	** ***	•		22		
		7.25	141	7.00	**	4 6	er .
" " imported 500 to 700 tons.							
Zine oxide 40,200 tons,	* *	4.00	111	4 93	**	**	11
" " imported in oil 16,000 " Dry, 2	St. t.	•••••	• • • •	4189-7			
interview to the total t	CM s. E4 12 1 20						
Plake graphite 1430 tons							
Flake graphite 1430 tons Amorphous graphite 2500 "	Value	5.50	141	4 98	**	**	44
Acheson's " 30 "		**!**	•••	0.7 x 889 C 7			
Imported graphites of all grades average	from	143 te	· 1	d tie	nes th	er alf	nounts
produced in America,							***************************************

nishing over one-half of the entire output of 14,200 to 14,500 short tons. Value \$0.50 to \$7.00 per ton.

Imported ocures, 7,700 to 8,000 tons. Value \$7.70 to \$7.90 per ton.

Spanish brown, principally from Maryland, 600 to 650 tons. Value \$17.70 to \$18.00 per ton.

Ochres of all grades were produced in 13 different States, Pennsylvania fur-

What proportion of these amounts were really applied for the preservation of metallic structures on shore or allow, it is difficult to determine; but one-fourth part may be taken as the yearly allowance to cover the effects of corresion in progress in some degree in about every metallic structure that meets the eye, and may be considered as the annual contribution to the coffers of corresion.

The greater part of the mixed pastes and paints of the day are adulterations, and are presented to the public in these forms the better to conceal the actual composition of the pigments and to cave oil; also to disguise the quality of the vehicle, as in the form of a paste or paint it requires chemical skill and time to analyze a sample of either. This, while applying in general to the mixed paint house colors, does not exempt large quantities of mixed paints sold exclusively as ferric protective coatings.

There are at the present day as pure brands of linesed oil, red and white lead, lampblack, and other pigments manufactured as any ever made. Possidy they are better on the average than those made one hundred years ago; but there are more that are a great deal power, and rendered more so by adulterations of the most barefaced character. There is a great advantage in the use of prepared paste, as the quality of the vehicle required to lang it to paint can be positively known, also the drives used, and what amount of these is necessary to meet any condition present at the time and place of applying the coating, details that in most cases cannot be known in advance or by the paint compounder, unless, as is too often the case, he makes but one kind, and fits at for the constant placed duty by the difference in price and the guilbelity of the

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spread a poor paint as a good one. On railway bridges, viaduets, and structural ironwork painted in situ, it costs for the painters' labor about twice the cost of the paint and in many cases four times as much, depending upon the character and amount of scaffolds or ladder-work.

This assumes that a reliable paint is used that costs about a dollar a gallon, that will cover from 300 to 400 square feet of surface for the first coat, and from 500 to 600 square feet for the second or a repainting coat. In the latter case the surface covered may be less, or the same as for the first cont, all depending upon the labor of seruping or the condition of the surface of the old coating, whether scraped or not. Obviously, the claim that a paint can coat 1000 square feet of surface or more, and prove as durable as those covering less surface, as above, is not sustained in practice, though it is always possible to get a doctored result with any paint, good or bad. Paintfilms that is, the oil rovering the atoms of the pigment are only from all to take inch in thickness, whatever the size of the pigmentatoms. It stands to reason that a thick coating of the vehicle will better protect the pigment-atom than a thin one. If the pigmentatom is susceptible in any degree to atmospheric influences, it will he less affected with a heavy conting of the vehicle than with a thin one. A thin coating usually implies that the oil has been reduced in density to render it easier to spread, and to be spread over a larger area, by the use of a larger quantity of solvents, either turpentine or benzine, than is necessary with any good quality of either raw or boiled linseed-oil.

Red-lead paint, from the large amount of oil in it and its great specific gravity, spreads over a large area, and it is these features that cause it to run or crawl on vertical or slightly inclined surfaces, particularly in the first coat.

A like result follows the use of flake-graphite pigments. The atoms of this variety of graphite, on account of their smooth surface and low coefficient of friction, appear to slide around in the vehicle before it dries enough to retain them in position when spread. The silien and barytes frequently mixed with such pigments to give a

cleaning, the plates still showed thin yellow rust-spots, that showed plainly, but of a darker color after conting with oil. The oil was scraped from some rust-spots under the oil on dry girders in the yard, and the yellow color of rust, so often found, was developed."

It is to be regretted that this engineer's views of what constitutes a thorough preparation of the ferric surface for its coat of paint is not an exception, but the rule in more than nine-tenths of the structural manufacturing establishments. Notwithstanding their claims to pre-eminence in their profession, they have yet to learn how to protect what they create; and that they are either incapable of this, or indifferent to it, the present condition of the ferric structures of the day is an unanswerable evidence.

If the superiors do not understand the importance of the proper preparation of the surface to be covered, or the character of the paint and manner of applying it, or give them the same or more consideration than they attach to other matters of construction, it will be next to impossible for the inspector or master painter to enforce good work. It requires a more determined stand on the part of those in charge of this branch to ensure good work, than in any other part of the construction details. Until the head officers are zealous enough to care something about the condition of the work after it has left the shop, and the men actually in charge of the painting are given to understand that they will have the unquestionable backing and support of their superiors in any stand they take against the present socalled practical methods of structural painting by unsempolous contractors, just so long will their work show their neglect in the rapid progress of corresion, that will not need scraping the surface of the conting to find.

The low grade of labor available for the painters' gang has much to do with the generally unsatisfactory results obtained. Painting can be slighted and still present a creditable surface that will pass inspection more easily than any other branch of hand labor connected with bridge or structural ironwork. Painting is as hard in muscular requirements as light blacksmithing or the vise-work of a machinist, of the pigment.

Probably the best results could be eletained if the man or firm who pays for the completed structural work appearated become their continuous importor to attend to this branch of the work with the distinct understanding that his orders were to be strictly enforced, and that his endorsement on the full rendered was necessary before payment of the same. This would ensure the proper properties of the amface, and secure enteful attention to the before mentioned necessaries, and

jie flieffitiff. it gittige, to eite be gare, efigioned o ein gann, Bingang, gann pon inginent. Intefif

These features are particularly apparent if red lead to the paint used for the shop cost, as any want of care in keeping it continually well stirred up in the paint-pet by the paidle stick (not by the brush) to prevent its "setting" is almost undetectable, and the want of care here governs the durability of all of the subsequent restings. The use of lampidack with red lead in a paint coating, while it delays the quick "setting" of the coating, does not present the rapid settling.

other conditions are favorable to service a first class result

he alone could be held responsible for the fixed tensity

engineers have proven to be better protected against correspon than the structures painted either by contract or by the most premiment of the construction firms, who, as a rule, are more anymous to get the work out of the shop, than for its future fate.

The pickling of structural ison with dilute arols to remove the mill scale, as done in some classes of ship and loader work, has sact with many objections. These objections are primarily the cost of the process compared with a rush cost of something demonstrated

railway bridges that have the several contrage of point applied under the direct appervision of one of the railway company's coverence of

paint.

When pickled and brushed clean of scale, the metal must be copisously washed in water and then dried if possible in the sum or artificially in a warm room or oven, and then whether machined or not, be coated with the first coat of paint. If a few hours classe before applying the coating, the surfaces will begin to acquire the thin blush coating of red rust, as described in Chapter III.

men century of rest rust, as described in Chapter III.

The use of the sand-blast at the final stage of the machining passes

and grim determination to apply, but the result in having a properly cleaned surface for the foundation of the protective coatings has been proven in hundreds of cases as the only sure method to reduce the maintenance expense of the structure. (See Chapter XXVIII, Sand-blast and Pickling.)

Many engineers are advocating the plan of having a coating of either boiled oil or paint applied to the iron or steel at the mill as soon as possible after it has left the rolls or hammer, and while the metal is hot. The hot part is the only part to commend. All metal as it leaves the rolls or hammer has a tough, thick or thin (as the case may be) coat of loose or partly loose scale that adheres for the time being, but on a short exposure to the air with a few changes in temperature, due to mill or storehouse conditions, releases its tension and is ready to fall off whenever handled, as in the course of loading and transportation. No amount of brushing that any mill employé would or could give to the metal in its hot or half-cold condition would remove this scale, and if the painter was present with his pot of oil or paint, it would get on over scales and all, and no ordinary inspector could prevent it, or be in any way sure that the contract requirements had been complied with in regard to the removal of the scale or the composition of the conting.

The mill coating is exposed during its application and drying to all the dirt, einders, and sulphurous gases of the mill, which are a fruitful cause of decay in a dried coating of paint, and find an easier field in the green one. The mill-coated work is not allowed time to dry before being loaded for transportation, which adds its quota of dirt and einders to the sticky paint.

All the subsequent machine operations are accompanied by more or less lubrication of the tool, and the oil used for this purpose is the cheapest to be had, and in general has been used over and over again; is dirty, sour, and more or less decomposed, and carries enough hydrocarbon to evaporate and dry down as a dirty surface skin, hard to distinguish from the coating applied at the mill. The sequence is that the inspector crowded to get the work out of the shop, and if at all carcless in the discharge of his duty, does not personally see that the scales, dirt, and machine-grease are properly removed. The

cedure at variance with what he thinks is a special function of his craft, hastens to get on the paint, and takes more excell to himself in being able to beat the inspector than to do a meritorious piece of work.

Rather let the material go from the mill or forge to the storeroom or construction shop, protected as far as possible from any unnecessary exposure to the elements. When machined, during which process the greater part of the mill-scale will be loosened up so as to be readily removed, and when the several parts are assembled in their relative positions ready to be riveted up for their permanent places in the structure, if it is to be done at the shop instead of in situ; then and there is the place for the inspector to determine if the several parts are not only properly machined, but also properly cleaned from the scale that has not been removed by the machining and handling. He should see that grease, dirt, and any remaining scale, tight or house, is removed in his presence, and the first coat of the paint applied in a manner to meet the atmospheric conditions at the time, and use a quality of paint that will ensure more than a guess at the future protective result.

Nothing can then serve as a cleak to hide the impactor's responsibility for the result. One inspector, and one impaction at the final stage, is better than a number of impactors and impactions strong over a claim of operations comprising months of time and hundreds of miles between the links.

Many engineers advocate the use of boiled oil alone for the first or priming cont, applied either at the rolling-mill to protect the metal during its transit from the mill to the construction shop, or at the shop when ready to ship for erection. The general reason assigned for this practice is, that the boiled oil "sanks into the scale and dries and prevents further tendency towards corresion."

This theory is absolutely without preed, from any standpoint. How far any oil or liquid can work into from it steel it the still harder mill-scale that forms on these metals, these Solons do not state. The way of such oil continue is in secural to correct more

the metal surface. There are hundreds of records of the painting of important railway structures, where the first coat of boiled-oil method was used, and, in the great majority of instances, the utter and rapid failure of the coating, and the extra corrosion of the structure, could be directly assigned to this so-called method of protection.

The weather-resisting power of an oil coating is almost nil compared with a paint, as before referred to in Dr. Dudley's experiments (Chapter I). If the advocates of oil coatings are so sure of its benefits as against a paint, why not make all the coatings of oil alone, no matter what it covers, a wire or an anchor? It will soak as far into one as the other. A paint coating can be applied as quickly and easily to any surface as an oil coat; will dry as quickly and as hard, and is in every way a better resistant to atmospheric or mechanical injuries.

A foundation coat of oil is a direct cause of the blistering and peeling of the coatings spread over it. It is seldom dried enough before the other paints are spread over it, to ensure a close adherence to the metal it covers. When the subsequent coats of paint are spread, the solvents and oils in them soften to some extent the underlying coat of oil, and a moderate heat from the sun causes the whole coating to blister or peel. Too much oil in a paint coating, particularly if the surplus oil is in or near the foundation coat, whether on a wooden or metallic surface, will generally cause peeling regardless of the pigment used in the coatings.

CHAPTER III.

1934 8 % .

Symbol, FE. Atomic weight, 56. Specific gravity, 7.77.

Iron is never found pure in nature. Its avoids for exygen is so great that it quickly forms terrous exide, bett, or the protoxide of iron. This also is never found free, and is obtained to obtain chemically pure, its affinity for exygen forming the sempeoxide. Fig. (and also the peroxide (the highest form of exide for any metal) in which two atoms of iron, and three of exygen are united, or 70 per cent of iron and 30 per cent of exygen.

In the latter form it is commercially known as iron exide or iron ore, and is found in all parts of the world in all stages of parity, and in combination with the exide of all the other metals in all proportions. The color of the protexide is a green his changing to a red-brown—that of the perexide is a blood-red.

An intermediate oxide the black numberic, left, three atoms of iron and four of oxygen - 72 4137 per cent of mon and 27,5863 per cent of oxygen is the purest oxide of iron.

The ferric anhydride, Lett, is not known in nature, but is supposed to be formed by fusing iron or its exple with nature. Its color is a deep crimson.

Iron at a temperature of 230°C, (446°E) accumbance freely with the atmospheric oxygen, becoming first covered with an extremely than film of magnetic oxide, Le₂C₁, of a light veltor veltor which gradually passes into red blue, and gray color. At a shifte heat, area forms in the air with a production of magnetic exide, the conduction being sustained for some time by directing a black of his agent the heated metal. At a temperature of 360°C (480°C), iron decomposes steam,

then in its freedom from sulphur. It hydrates on exposure to the air or moisture to Fe₂O₃+ Π_2 O, and can be reduced to metallic iron the same as any iron ore.

The precipitate formed from metallic iron when corroded under water is the sesquioxide or peroxide of iron, Fe₂O₃, plus three parts or 24 per cent of water, and is red rust, Fe₂O₃ + 3H₂O. It is a dull reddish-brown color, nearly a pure oxide, containing only such other metallic oxides as the iron contained from which it was corroded. It is comparatively free from sulphur, more so than the best hematite ore.

Oxides of Iron.

If purity of an iron-exide pigment is any factor to prevent corrosion, these pure exides ought to be better than any iron-exide ores; but they are not, and plainly show that the failure of all iron exide-pigments to prevent corrosion on a ferric body, or to add any resistance to the decay of the paint coating, lies in the natural inadequacy of a ferric pigment to resist its own inherent weakness, namely, conveying excessive amounts of exygen with a tendency to excite electrolytic action.

Experiments determine that bright iron placed in an atmosphere of dry oxygen, or of dry carbonic acid, will not rust; when put in a damp atmosphere of oxygen, it rusted slightly; in a damp atmosphere of carbonic acid, a small quantity of white carbonate of iron is formed on the surface of the bright metal, but no rusting takes place. When, however, bright iron is placed in a damp mixture of the two gases oxygen and carbonic acid it is rapidly oxidized into copious excrescences of red rust.

In the opposite direction, to prevent rusting, a strong solution of earlianate of soda preserved needles and steel instruments, bright and untarnished, after thirty years of exposure, and would probably do so forever

Bright steel or iron objects remain untarnished in an atmosphere of dry muriate of lime, also in the dry carbonate of lime. Iron immersed in lime-water, caustic potash, and caustic soda does not rust; though the lyes absorb carbonic acid, they do not absorb oxygen.

measurably thick and prevents any further exclusion. On the contrary, iron-rust once formed on a ferric surface never ceases its action so long as it is in contact with it. Rust produces rust

The blush of oxide that appears upon the surface of a piece of bright, clean iron, such as is left from the action of the sand blast or of a grindstone forming after a few hours of atmospheric exposure and can be wiped off by the hand; or the piece of red rust from the iron-scrap heap; or the scales from the bottom or frame of an iron ship; one and all are the peroxide of iron, Ec. O., plue three parts or 24 per cent of water. When calcined to drive off the water, they

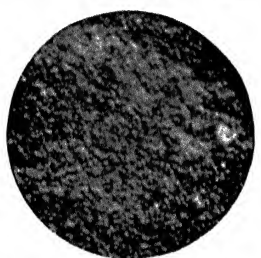


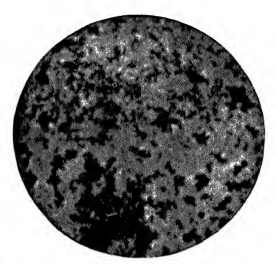
Fig. 4.—Rust produced on a clean relied-iron plate exponent to atmospheric influences for 20 days, a Andre

become precisely the same ferric oxide contained in any iron ore, and will reduce to metallic iron or grind to a pigment the same as any iron ore, however they may be designated or juggled with trademannes.

In the corresion of iron from any cause, for every 8 grains in weight gained by the iron, 46 cubic inches of hydrogen weighing 1 grain are set free. With each cunce of gain in the weight of the iron, 2515.625 cubic inches of hydrogen (-1.4555 cubic fact) are

titaniferous sand, found on the seacoast in many parts of the world, exposed to sea-water and other sources of oxidation and friction, has remained unchanged for thousands of years. Its use for a pigment is not satisfactory, on account of its black color and the difficulty of grinding it.

Specular iron ore is also but little affected by oxidation, and is a nearer approach to a definite compound of iron and carbon than any other known ferric substance: iron, 94.85 per cent; carbon, combined



Pto. 5.—Rusting of a clean rolled-iron plate from a single application of water and left to dry. (Andes.)

and graphitic, 3.50 per cent; silica, manganese, sulphur, and phosphorus, 1.65 per cent. It is an anhydrous ferric oxide, found in Nova Scotia, in the Isle of Sicily, and other parts of Europe, where mines of it have been worked for 3000 years. Its black color and hardness prevent its use as a pigment, though its resistance to corrosion is almost equal to that of the Bower-Barff surface.

The clay-iron ores from the coal measures, the spathic, bog, and many other iron ores, contain a very small amount of iron, and so large an amount of silica and other mineral substances that they are The red and brown bematite non-ories, composed of the sequioxide of iron. 70 per cent metallic non-ined 30 per cent extrem, plus
water, plus variable percentages of immetal substances, plus carbonic,
sulphuric, and phosphotic acids (see following analyses) are the
principal metallurgical non-ories, also these need for the production
of pigments under the name of non-oxides, Left,. This obsticial
symbol, name, and product is subject to more commercial pigglery
to meet trade requirements than any other pigment in over 1t is
made to cover all sorts of combinations discree in composition and
character, supplemented still further by quantities of socialled mert
bodies, more unstable than these in the ore with which they have
been brought into forced relation.

These hydrated ones when cabined to expel the moisture outplinite, phosphore, earlience, and either erroments supposed cauly evaporated neits become the anhielders or supposed muttal, lead, plus about 2 per cent of water in a combined form, plus the and elements that frequently amount to 2 per cent. Only the following cents are dispersed in the comparatively low heat of the masting process, and are not aboild consumed in the high heat of the blast furnace, as manufacturers of metallic iron find to their amountaine.

The lime, magnesia, alumina, silva, manganese, and other exples, whether in a combined state in the ore, or as added free substances at the time of reasting by calcination, become caustic and his greecopic, and when ground to a pigment form, absorb moisting from the atmosphere, slack, changing their character again more or less to a floccus lent or a powdered state. They do not beaut in the slightest degree to the oxide of iron or base, are no more connected with it or to each other except in a haplaneard arrangement of their disrupted, separate natures—than the same substances would be if collected from a sand-bank.

No mechanical process connected with their investigation into

catalytic power of nearly all finely powdered substances to condense moisture and gases from the atmosphere, which the porous nature of the paint coating readily absorbs. If sulphur is present in either the pigment or vehicle in any recognizable quantity (as it nearly always is), it furnishes an additional excitant for the electrolytic action. This electrolytic action is further intensified by the unequal composition of all iron ores, whether rousted or not. The process of roasting—always an uncertain one—does not affect the ore equally. Lumps improperly roasted, or from their composition affected differently by the process, are difficult to detect in the hasty and generally poor assorting or picking-over the ore receives before pulverizing. The same uncertainty in the composition and assorting attends the unroasted ores.

In the pulverizing process there are many larger and harder particles of the ore that would not pass a No. 50 mesh sieve, if the pigment were bolted (as it seldom is), and would much less pass a 100 mesh, to which size all pigments should be reduced.

The finer the pigment the more thoroughly will it incorporate with the vehicle and protect itself and the surface covered. The destruction of any particle of the pigment will not render the coating so porous as when a larger atom is removed to permit access for the atmospheric moisture and gases. These lumps act as centres to determine the corrosive action, and in a measure explain the erratic action of all iron-oxide coatings. In nearly all rust-spots, one or more of these hard particles will be found, and particularly so wherever pitting has commenced.

The brown hematite ores are claimed to be practically free from sulphur, therefore the best for a pigment; but the best brands of this variety of ore prepared by any one of the many manufacturers of unroasted iron-oxide pigments have not proved to be in the slightest degree any more reliable in composition, or any better protection against corrosion—whether used as a straight paint or mixed with adulterants—than those prepared from the red hematites. (See

The dirty purplish-brown or lifeless color of the brown hematites, even when freshly applied and aided by the gloss of oil, is not agreen-

following analyses of both pigments as used in commercial paints.)

out

protective agent, and the low cost of the whole line of iron-oxide pigments, are the great inducements for their production and use for ferric contings.

ANALYSES OF HON ORDS.

Hr Vanmen Assesses

Committee granute from Add to 1 % ?

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There is a wide difference among these comparatively few and better quality ores, selected from many hundreds of ore-bods, on account of their purity and high percentage of metallic iron.

The second secon

Many other mines furnish ores that are worked for the other metallic and chemical substances they contain, as nearly all the other metals are found associated with iron. All iron mines are noted for the variable quality of the ore taken from the same or from the adjoining hel, or from different parts of the same vem in each name. The hematites are not exempt from this feature, whether used metalagainst corrosion than those containing 80 to 95 per cent of the sesquioxide. This will be apparent by reference to the composition of the iron-oxide pigments given in the tests of commercial paints, Chapter XXX.

Iron-oxide Pigments.

The red hematites furnish a brighter-colored oxide than the brown hematites, whether roasted or not. The small amount of sulphur in the red ore develops in the process of roasting the dull-red color into a brighter red, simulating the Venetian and Indian reds so desirable to produce, but not always possible to get without doctoring the furnace product subsequently with substances more complex and unstable than the iron oxide itself. (See inert pigments, Chapter XVIII.)

The rousting process is a sensitive one. A few degrees of higher or lower temperature, or a little difference in the period of exposing the ore to it, or in the manner of cooling down the furnace, cause a great range in the color. The more sulphur in the ore the brighter the color.

There are from 10 to 20 per cent of moisture and carbonic neid in all iron ores as they come from the mines. If these are not driven off by roasting, they will not be dissipated in the pulverizing, and will be carried by the pigment into the mixed paint to its detriment. The use of an uncalcined iron-ore pigment is a long step toward an early corrosion of the ferric body over which it is spread.

The following analysis of an iron-oxide pigment made from a special red hematite roasted ore, one of the oldest and best known of this class of pigments, and the use of which as a special brand is probably greater than all of the other brands of iron-oxide pigments in the world, is of interest for comparison with an unroasted ore pigment:

MANUFACTURERS' ANALYSIS,

l'eroxide of fron	٠,	,						,		,	,						Da.	11	IM.I.	rent.
(Equivalent	ì	11	11	10	ŧ:	۱۱	lie	•	ir	¥ 8	11	,	31	Ì,	1,	ii	•		•	
per cent.)																			
Silien combined.								,			,	,				,	-16	03	44	11
Lime																				**
Moisture			,					,				,					1.	59	**	**
*															•	•	40	** *	4.4	

either wood or iron surfaces, if there is any protective value in nonoxide. However, notwithstanding its almost uncontested use for over thirty years, on account of its low cost, agreeable color, and much landed protective virtues, it proved as uncontribution to both wood and iron contings that various railway companies—the largest consumers of paints shave reduced the 50 per cent of this perseade of iron admissible in their inveducator paints to 25 per cent. But this change has not resulted in any marked improvement in the protective qualities of the paint when applied to ferric bolics, nor are better results apparent upon usealers surfaces

The following analysis is of a bown beneatite uncertaint on one, With a number of other branchs of similar composition, it has been largely used by construction and railway engineers upon hundreds of the most important ferric structures in the country, whose serious corrosion, after but a short period of exposure, led to a special examination and report on their condition to the engineering firms responsible for their erection and condition

Peroxide of iron	** 1	43 1	\$ PW : \$	9"#"12 <u>#</u>
(Papirolemat in montaline seems,				
65,128 per cent :				
Hillen combined	. n	3 34	* *	14
Alumius constituet.	**	. 3 % . 4	. 7	. 1
Line and magnesia	8 9	1343		"
Organic and volutile	黄素	1 4	**	**
Sulphuric nekt.	4 4	441		**
Moisture and less.	3 %	1 %		* 4
	1,000	1 51 1		

This is a high-grade metallic moneyer compensatively dyes from only plans, and whose metit as an units correspond pageness was greatly commended (by the manufacturers) for a straight paint free from the usual class of met adulterants. But its protective results as detailed in the said report were not better than the other adulterant objective pigments, or the other resatings of mixed or unknown acommonition.

The following is an analysis of an iron while pignoest that is

Carbonate of lime	30.19	"	"
Sulphate of lime	14.05	"	"
Clay and silica	19,90	"	+ (
Alumina	8.48	11	11
Magnesia	0.52	"	"
Water and organic matter	0.44	* *	"
ân â	(A) 2-m		
1	(00,00		**

Additional examples of iron-oxide paints and their erratic action—both mixed and straight pigments—will be found in the article on paint tests, Chapter NXX.

It is claimed that iron-oxide pigments, being the peroxide of iron, are incapable of further oxidation, and when ground with the vehicle are indestructible, and their capacity to condense atmospheric moisture and gases ceases. This is true as long as the thin film of the dried vehicle only $\frac{1}{2} h_B$ to $\frac{1}{4} h_B$ inch in thickness—remains in place on the external surface of the pigment atom, and no longer. The same causes that remove this film will affect the other part of the vehicle, in which the pigment atoms are imbedded. The vehicle, passive of itself to condense atmospheric moisture or gases, is porous and absorbent; and passes them on to the point where their decomposing action can take effect, if not on the iron-oxide atom, then upon the less resisting mineral substances associated with it as a pigment.

With the possible exception of silica and barytes, all of the so-called inert substances, usually mixed with iron-oxide pigments, are porous and absorbent of the vehicle and gases that reach them. The protection that these inert substances receive from the oil is no greater than the oil affords the iron oxide atom, if not less, owing to the unreliable character of their composition naturally. If they have been mixed with the iron ore during the process of roasting, they are rendered more unstable, and readily pass to a lower plane of resistance, as mentioned before.

It may be questioned whether iron oxide is incapable of condensing moisture or gases. It induces and promotes oxidation in and other oils is to absorb the glyceride element that, unabsorbed or unchanged, in all fatty oils delays the drying process, condensing the atmospheric moisture and gases that act below the external film of the drying oil, thus laying the foundation for a blister with subsequent corresion of the coated surface.

The power of iron exide to absorb the glyceride is about twothirds that of red lead. If the non-exide atom is incomplete to the presence of sulphur that may be presented to it in any form, the other associated mineral substances and schule are not, and a very small percentage of any and will set in motion the electrolytic action so fatal to ferric substances.

As a class, the inert pigments are electro-positive to the irinuxide atom, and are the first to be affected by any electrolytic action inaugurated by their presence in the panel.

The iron exide atoms are electronegative to the ferry surface over which they are spread

In iron saide and sinc cable maximes, the mon atom is electronegative to the sinc atoms, which are spinish destroyed. If any engine or enquer wade is present in an iron caule payment, the mon calde is electro-positive to the enquer and is destroyed.

Mallet's experiments determined that respect and size in any form, added to, or in contact with non-in-any form, increased the correspond a covered iron surface fit per cent in a given time, requer, without the sine element. It per cent.

The irregularity of the distribution of the atoms in a compound irrateoxide conting their difference in one and character determine the points of corresponds, which once established and only with the complete failure of the coating.

From five to ten per cent of the sulphate of line (Catelly) is generally found in iron oxide paints or pigments. It is aimally speciallied by the consumer that it shall be fully hydrated, or that it shall contain not more than one part of water. The effect of the minimize is to aid any sulphur element present to communical presents are work of disintegrating the conting. From its war at any other paper.

carraces. The brown exides stand the best upon ferric bodies. The Venetian red exide, from the old iron-exide mines, had a peculiar preservative action on the surface of wood. Two or three coats from the pure materials have outlasted the record of their application and the lives of the painters that spread them. These exides and white lead form a hard mastic covering, and miless spread over misensoned or wet wooden surfaces, are not liable to blister or ped.

Many of the irreconcilable discrepancies in the use of iron-oxide paints can be attributed to the careless method of preparing them. In general practice, it is never ground with the oil, and but seldom machine-mixed. The dry pigment in the ratio of six to seven pounds and about the same weight of oil (or three fourths of a gullon) are placed together in a tub, and after a few hours of sooking are simply stirred up and spread. If any large quantity of paint is so prepared, it is almost impossible to secure thorough incorporation of the pigment and the oil, owing to the different specific gravities of the several substances composing the pigment, which vary from 2.2 to 4.9. This manner of mixing is strongly recommended by the iron-oxide trade to secure its use, at the expense of the life and effectiveness of their product, which many times might be more creditable were better care taken to render it deserving.

The longer that iron-oxide paints are ground in the full quantity of oil they require to form a *paint*, the more lasting they will be, and this effect is equally apparent in all pigments.

The unsatisfactory results due to carcless mixing are aggravated by the use of large flat brushes that act as mops to carry or slap on a large quantity of paint, imalequately, in this way, brushed out. Such brushes carry air into the conting, rendering it more porous in drying than it otherwise would be were heavy, long-bristle, round brushes employed. The same objections exist where the conting is applied by the air-brush or spray apparatus, only in a more marked degree. See Chapter XXXI.

There are many tests for the adulteration in iron-oxide pigments or paints of too extended detail to be entered into here. A ready ing the use of from exide pagments. Ferschus (1858) mentions that on houses in Sweden, painted with from exide three hindred years ago, the conting was still in fair condition, and the west work well preserved. The western houses and workshops in many parts of the United States bear testimons equally in faxor of this class of paints, in the general good condition of the paint and structure after exposures, without repainting, of fifty to eights years. But in every case the cause of these excellent and exceptional results was in the use of better materials and better methods of application than is the present-day practice.

Paint-tradeliterature frequently extend that the time roof of Independence Hall in Philadelphia has been protected for the past one hundred



Fig. 6,.... Correspond of the present day make of the plate after stronge in a device find for ten years. (Andre)

and thirty years with an iron cylde paint, giving this as an unanowersble argument for the protective character of these pigments applied to metal surfaces. The facts in this case show that the plates with which this particular roof was laid, as well as many others at that period, were double-control with pure block tin, free from lead, sincand no amount or kind of paint spread upon an interior quality of tin-roofing metal can prevent this internal corrosion, though it may coneeal its presence and progress, and possibly fill up some small holes in the early stages of the decay.

There are scores of tin roofs covering important buildings in the Canadian Provinces that have been laid for nearly a century, as bright and uncorroded now as when first laid, never having had a coat of paint to protect them.

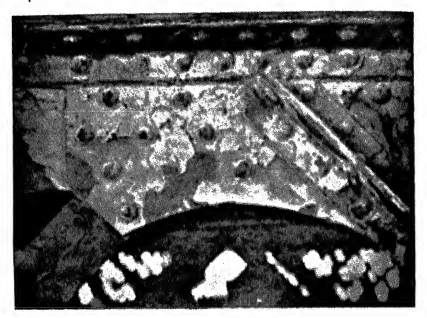
Pure block tin is unaffected by atmospheric conditions, almost as much so as copper. It was only when poorly cleaned plates, poor tin for the coating, acid flux instead of resin for soldering, and careless methods of laying the roof generally, came into vogue, that we began to hear of the virtues and need of an iron-oxide paint to prevent the corrosion of a tin roof. Granted that a good quality of tin roofing is none the worse for a coating of paint applied a year or two after the roofing is laid, yet it is quite as essential for the future life of the roof that the paint should be also of good quality.

The original mines from which the iron-oxide pigments known as Indian and Venetian reds were taken have long been exhausted. These old mine pigments required no roasting or doctoring to develop their color, or to correct any acid elements in them. The reputation of these pioneer oxide pigments, like that of the "Old Dutch Process" white lead, has been assumed to reach and cover the advent of scores of substances bearing little resemblance to their progenitors, except in name, and even this is not exempt from the greed of some modern paint compounders, as the many prefixes and trade-marks bear witness.

Fig. 7 shows the protective character of an iron-oxide paint applied to a railway viaduct not properly cleaned from mill-scale before painting, and when painted was exposed to combustion gases, einders, dust, and moisture.

Prepared from-oxide paints are often brightened by the use of aniline colors, but are not durable. Burning a sample of such paints over an alcohol lamp will destroy the aniline, and leave the iron oxide its natural color, expessing the cheat. The tendency of all iron-

power. The clays also alsorb water and harten the decay of the paint.



F182. 7

Copperus Oxide.

The chemical composition of cooperas is Fe 50, 711,0 - one part of iron plus one part sulphanic acid, plus seven parts of mater

Copperss, the waste product of many manufacturing processes, is largely used to produce an exide of iron pigment to reacting the crystals. Six parts of the water are driven off to: a heat of 114° F. but one atom is still retained at 250° F. At a red heat of decomposes, giving off one part of sulphurous exide, leaving a large ferric sulphute, Fe₂O₂SSO₂; and, more strongly heated, it leaves a pure ferric exide known as Coleothar vitrial

As usually roasted for an iron exide adgment, from three to five pounds of termalla, lime, or chalk, to one of the ore, are just in the

process is a sensitive one, the color of the product being the brightred color pigments. Venetian and Indian reds. Both colors are due to the degree of heat employed, the length of exposure to this heat, the manipulation, manner and time taken to cool the mass, etc.

All of the sulphur is not dissipated by the heat nor absorbed by the line in its change from v carbonate to a sulphate. The line changed to gypsum or left free, being in great excess of the amount that is allowable in any pigment, is removed to some extent by sifting, or other means, before grinding the furnace product.

Copperas oxide requires great care in its use, either by itself or mixed with the dead color iron oxides or other pigments to bring up their color, as the sulphur goes into the paint with the usual results. No amount of free adulterants or inert substances have any material effect in neutralizing it.

Copperas ferric salt (a protosulphate of iron), Coquimbite, is found native as a hydrate, containing uine atoms of water, Fe₂O₃BSO₃ + 9H₂O₃. It occurs in layers several feet thick in fine grained six-sided pyramid crystals. Its preparation for a pigment is similar to that described above, and the resulting disintegrating effect in the paint is not measurably different.

Yellow Ochre.

Other is a hydrated oxide of iron of a strong yellow or brownyellow color, generally containing less than 40 per cent of iron oxide. The others are among the oldest of pigments. Samples have been obtained from Pompeli in all stages of preparation from the ore to the mixed paint. They were used in Greece in the time of Pliny, and in Old Egypt. They are the most stable of the yellow colors, and are the principal pigment in the present freight car colors.

Ochres are clays tinted with the oxide of iron and manganese, and hygroscopic in character, carrying from five to fifteen per cent of water. Dried artificially to expel the water, they change color to pink or red the same as all other iron oxide substances.

Their vellow color is chiefly due to the iron oxide, and the more

The covering power of the ochies depends upon the amount of lime or chalk in them, which reduces the coloring power by rendering them translucent. They require from sixty to eighty per cent of oil to form a paste, and the added quantity of oil to make them spread makes them slow divers. They blacken a little in time exposed to similable, but the change in tone is evidently more from the darkening of the oil than from a change in the pigment.

The best brands of other are the I reach;

Composed of clay.... 69 5 to 73 8 per cent, Oxide of iron and manganese 23 5 ' 25 0 '' Water.... 7 0 '' 9 5 ''

French ochre has a large spreading power, as it absorbs a large quantity of oil, and it holds well to wooden surfaces. It should be ground in raw insected, and if a thinner is required, raw oil should be used. White ochre has the property of hobbing well to wooden surfaces from the large amount of oil taken up to it, but does not bond well to any overlying coat of white lead, and tends to east it off by pecling. This action can be associated by using a small percentage of white lead in the priming schire coat.

The English Oxford and stone others are among the best brands. The Roman others grade with the best Havre, while the lower grades of French others are poor and possibly lower in covering power than the best Bermuda (Virginia) or other American brands

The name of an other signifies, like all other point names, little, unless the material is furnished from a responsible because from Even these cheap earths have to hear a share in the general burden of adulteration that is the order of the day, by an added quantity of clay, chalk, and barytes (the latter to give weight), but all inpute the covering power of the other. Their presence is usually denoted by the increased amount of oil required to bring the dry other to a paste.

There are many mixtures of other as the basic pagment for a

weather requires the least oil.

The better brands of ochre as the basic pigment for freight-car colors form very durable paint coatings, whose life is generally equal to that of the car. The cheaper grades were formerly used to a great extent as cheap paints for tin roofs, but the large amount of free sand, lime, and other uncombined mineral substances, acids, and moisture that they contained, with the coarse way they were calcined and ground, rendered the contings short-lived and unsatisfactory. They required a large amount of oil to spread them, even with a white-wash brush. They dried or lardened rigidly, did not bond to the tin, and the rate of expansion and contraction from the action of the sam was so materially different from the tin they covered that

Umber.

they soon eracked, blistered, and flaked off.

Umber is an argillaceous brown hematite ore, essentially $2\text{Fe}_2\text{O}_3\text{SiO}_2\text{H}_2\text{O}$, with alumina and manganic acid. Specific gravity 2.2. Originally obtained from Umbria, Italy, now chiefly from Cyprus.

As a pigment it is used in both its raw or natural state, and when calcined is known as burnt umber. When calcined at a low heat it turns a dark brown; a stronger heat dehydrates it, turning it to a red brown and softening it. As a ferric paint it has no special quality other than the ochres. The cheapness of the pigment is more than offset by the amount of oil it requires for a good spreading paint.

I inher is used as a drier in boiling linseed oil, and furnishes an oil of good color; but unless used in large quantities, does not make so rapid a drying oil as the lead, zinc, or unuganese driers.

Spanish Brown.

Spanish brown, an iron oxide or other, containing thirty to fifty per cent of clay, is inferior in color and covering power to umber, but is of lasting value for a roofing paint, as the clay, which has at all times a strong affinity for moisture, will, when properly calcined, take up seventy to eighty per cent of oil, and this oil, protected from well protected as on the horizontal surface, will absorb moisture, soon corroding the ferric surface. If two or more ochre contings are spread over one another, the last conting is hable to peel, hence for tin or metal covered roofs, one heavy cont to better than two or more thinner, unless the latter are applied after an interval of a year or more. The more clastic the first conting of this jugicent, the more durable; but the greater will be the tembersy is east off or crack the second of other coats.

CHAPTER IV.

RED LEAD.

Metallic Lead. Symbol, Ph. Atomic Weight, 206,9, Specific gravity, pure 11,145; connervial 11,335 to 11,388,

LEAD occupies an important part in the arts and manufactures of the day, and requires a greater range of chemical and mechanical processes for its production as a pigment and more care in preparing and applying it for a paint than any other pigment.

In its mineral form it is associated with all of the noble metals, also with copper, tin, zinc, bismuth, antimony, arsenic, etc. Some of the baser metals are always present in commercial pig lead, and affect the character of the pigments prepared from it by the processes of calcination, exidation, sublimation, corrosion, and precipitation.

There are twenty ores of the metal known to the mineralogist, but metallic lead is produced from the five following minerals, the analyses of which indicate not only the character of the ore but also of the metallic lead and pigments made from them:

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Metallic lead forms five exides that in one or more forms are the result of the heat or chemical changes produced in the metal in converting it into a pigment. They are:

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Red Lend, or Minium, is the principal pignoest produced from the oxides, its specific gravity being 8.5 to 8.91, depending upon the purity of the lead "Massicot" or lithurge, from which it is made. The color also depends upon this point, but in a greater degree upon the temperature employed in the explanion of the material, the uniformity of the heat, the manipulation of the material in the furnice, the length of time exposed to the heat, and the rate and manner of cooling down the furnace and its contents

Special furnaces and processes are required in its proposation that differ materially in the various countries where red lead is produced, also in different manufactories, and there is a difference in the materials employed.

Briefly described, these processes are. First, by the cupellation furnace which converts metallic lead into lithrage in about 21 hours. Second, a reverberatory furnace in which lithrage is converted into red lead. Third a reverberatory furnace in executive that reduces metallic.

this mass when broken up is in the form of thin yellow or brown scales, and in this state is known as flake or glass-makers' litharge. This, when ground in water and dried, changes its color to a buff, and is the ordinary commercial litharge.

When the crude lithurge powder is again moderately heated in a reverberatory furnace or oven and exposed to a current of air with continuous stirring for from 26 to 48 hours, or until a sample drawn at a low red heat appears of a dark-red color turning to a bright red on cooling, the furnace is closed and allowed to cool slowly; a condition most essential to success in the color, that if not satisfactory, requires a reheating and cooling of the product now known as red lead.

Red lead made from lithurge (from the imperfect oxidation of the lithurge) contains a larger amount of the protoxide of lead than that made from the earbonate or white lead, where, on account of the finer condition of the material, the oxidation is more complete, more quickly effected, and generally of a better color and quality.

With this complex chain of operations there are many trade secrets to secure not only a uniform quality of the pigment, but its color. The latter nearly always remains an uncertain element even with the best of attention given during the whole process.

Red lead is found native in many localities mixed with the other ores of lead, probably resulting from their oxidation by natural causes. Chromate of lead (Ph CrO_b). Specific gravity, 4 th to 5.2), the neutral, or meta-chromate, known as crossiste or lehonante, is a native red-lead ore found in commercial quantities in many parts of the world. It is in the form of translucent crystals of a vellow color with various shades of other colors, and is associated with decomposed gueiss and granite. The method of converting these red lead ores into pigments need not be described here. All of the vellow and red chromates of lead are obtained from crocosite. They are strong colors, and do not decompose on exposure to the air or light.

Red lead to one of the heaviest and most expensive pigments, also the most difficult to prepare for a paint or to spread. It is more superptible to adulteration, and is more adulterated by interests initial

able to its improper preparation and application, and where the failure should have been foreseen by the engineer or master painter, and where carelessness, indifference, or ignorance of the conditions to which the conting was to be subjected, were the jame factors of its non-success.

When obtained from a reputable manufacturer and properly prepared with a suitable vehicle and spread under known conditions of its future service, it has proved to be one of the most reliable nigments. Its color is distinctive, hence it is not favorable to the use of adulterants. Buck dust, more exide and leavates time design its high color, but are detrimental in all other respects. Chalk, gypanu, and other light color and low specific grasits substances are often added to consent the tendence of sed lead point to "bet" in the paint put while applying if, or to present its "except" when spread upon vertical or inclined surfaces. All such adulterates are easily detected; they do not prevent the set or crawl of the paint, and are the principal cause of the fathure of the coating. For the foundation cost upon ferre bodies, it will cover about as much hunface as any other paint applied under the same conditions and with the same effort on the part of the painter to brigh it out. The latter factor in frequently time much to stanger made no wath white a white wash

As a first coat on ferric bedien, applied at the mortables, its index shows at once any material injury to the counting due to the usual handling and transportation, also readily indicates if the greate and dirt due to the machining processes in the slope or received during transportation and crection have been properly attended to or not It is a prime watch dog in this respect.

Its tendency to settle in the paint pot also "to set" and the necessity for constantly stirring it up by the painter probably lessens by a small amount the number of square feet of surface he can spread in a day. This is probably an adjection to its use that is effect by the many points in its favor. The rapid set of red lead when mixed a peculiarity of this pigment, is no objectionable in a paint before

and will not again bond the sand or to the masonry. The red lead will not recover its combining power that ensures the mutual bond between the atoms of the pigment and to the surface covered, be it wood or metal. When in this condition, if it is applied to a metallic surface, it "crawls," as it is called, and presents an appearance more like that of curdled milk than a paint, and the actual protection of the body covered is due to the vehicle only. It will add but little to the covering power other than what any adulterating substance would do.

The setting of red lend is due to two chemical reactions, namely, a combination between the litharge of the red lead and the glycerine element in the oil; also a combination between the fatty acids of the oil and the litharge, forming a lead soap, quite a firm substance, but one not favorable to the durability of the paint.

Many of the failures of red lead coatings, if rigidly traced to their source, would no doubt be found to have been caused by the care-lessness of the painter in not keeping the paint well stirred during its application, or in preparing too large a quantity for immediate use, or by using the paint left over from day to day, or from another job.

Red lead and Lampblack Mixtures to Delay "Setting."

Iron oxide, zine oxide, barytes, gypsum, etc., added to red lead to prevent "setting," are objectionable and ineffective, as before noted, Lampblack from 1 to 1 ounce per pound of red lead delays the setting action and enables the red lead to be prepared as a paste to be used in the immediate future, when it is thinned by additional oil at the time and place of using. The bright red of the minimum is modified by the lampddack to a chocodate codor that may be light or dark according to the quantity of the lampddack used. Lampddack of itself is an excellent pigment, is electrically passive or neutral to all pigments; and by natural formation is so finely divided that it mixes easily and thoroughly with the oil and red lead without deteriorating the quality of either

hardening by simply agitating or rolling the barrels daily. In one case the paint barrels were left inclisturised for four months, and though the red lead had settled in some degree, when it was stirred again the paint spread as well and dried as finally as though freshly mixed.

Lamphlack containing sulphur in any approximate amount should never be mixed with red lead for a conting. Coronicl most from chimpeys or furnace-flux, ground latinimous coal or color, or the most from most of the petroleum or heavy numeral oils contain oulphus enough to cause the prompt failure of a red lead conting even when used to the amount of 1 of an ounce per pointed of red lead.

Mr. Ball, master painter Pennsylvania Radioad, 1897, reported the result of some of his experiments with "productive paints for metallic parts of cars and trucks," via

First. Hed lend and raw liment oil, with litharge as a dract.

Secretard, Read lead and rise lineared soil, not struct

Maigel, Raml langel jangel flangungebeltaunde, umgenial gelanden

Pourth, Red lead one part, langulant three parts.

Pafele. Margeregel bernande geral nangeler auf granes.

Sixtle Mexican graphate

All paints were mixed from the same quality of raw linearhold; none but the first had any direct

At the end of lifteen months of atmospheric exposure at the car shops their condition was as follows:

Namesheer Chair land finited in the our thream ogwith

- "Two was intact and appeared to be in combition to recent
- "Here had sometal all as a decided or of places, observing that.
- frante with its in will wonders are in latter the
- 4. Pieres Mitte anenniegengengenge fürenne.
- We have in perfect condition, the same as the straight red-

The law against the second sec

ration leaves the coating more porous than it would be if the paint dried naturally; and it deadens the gloss. Japan driers for red-lead paints are less objectionable than the turpentine. They are heavier and add resinous matter to the paint in drying that is of the same character as the vehicle with which the red lead is ground. The drying of japan is principally by resinification and not wholly by evaporation, as is the case with turpentine; with the added japan the period of drying can be governed as required by using a small proportion of raw hisseed oil.

Excerpts from a trade catalogue * relative to the use of turpen-



Fig. 8 Dried film of red lead energeted with rast's limi. Taken from a painted attacture who holowed no rust on outline. (M. Loch).

time and ted lead way: "We are not prepared to advise the use of turpentine in chips ards, where, owing to time contracts, it is often necessary to paint in damp or freezing weather, though the practice in the Marine Department of the Maryland Steel Co. from many years' experience in the use of red lead is, viz.: Use three parts of linseed oil (raw, presumably) and one part of turpentine for the first and second coats, with sufficient drier (presumably inpan) to set

"Theoretically, and for dry, bught weather, red lead should be used with raw insected and need need red lead itself being a natural drier. By so doing, the chemical union between the parameter and oil is most complete and the resultant point to most chirable. However, for ordinary service add a small quantity of jupan or use boiled linseed oil. By so doing a more viscous vehicle is had, which better matring the heavy particles of the red lead, thereby preventing the running on vertical surfaces, and possibly giving greater covering."

Red lend Pand Mastures.

Many railway engineers favor the following mixture of red-lead paint for their structures where the time of deving to of little moment, For the first or priming roat, 20 points of red lead and I gallon of raw linearded. No driers for the second and third roats a paste made from 60 points of hydrated subplace of line, 30 points of lamps black, 5 points of red lead, making 100 points of pigments, to which 20 gallons of boiled linearded are added, making 30 gallons of paint. This makes a fair drying paint of a dark or dirts graves become oder, weighing about 84 points part of a dark or dirts graves become root to peant to prevent corresion lies in the red lead and lampslach, the subplace of lime adds the stuffing for quantity without contributing anything of a protective character to the mixture, and the resering power is pass.

If any of the low-carbon amorphous graphites were substituted for the sulphate of lime in this formula, the point would be better in all respects for conting ferric bodies.

Mulder's Experiments with Cheap Red lead Points

Mulder, in his experiments to produce a cheap point, smell builted linear-built containing 25 per cent of red lead. He smed is parts of this oil with every one of the following maximum. The iron exide used was Cartier's (Belginus, that analysed as follows:

of some merit.

Red lend 25 per cent, iron oxide 40 per cent, gave a very good.

conting.

20, 40, 60, parts of red lead to 100 parts of iron oxide gave excellent results. 20 to 90 parts of red lead to 50 parts of pulverized red roofing tiles gave a thick heavy coating.

40 parts of red lead and 100 parts of pulverized red roofing tiles gave an excellent coating.

20 to 90 parts of red lead and 100 parts of pulverized ironstone gave a paint of distinguished excellence.



Fro. 9 Photonic regraph - two of a film of red had paint showing grains of surious sizes, all mark or less ones sted with air buildies. (M. Toch.)

Other mixtures of red lead favorably reported upon by experimenters are red lead, zine oxide, and Blanc-Fixe. The latter substance serves to hold up the red lead in the mixture during the first stage of drying and prevents its "ercep." Amorphous graphite is also used instead of the Blanc Lixe for the same purpose. None of these substances or any others employed for this purpose can be

The bright lustre of red lead is often toned down by Venetian red. This pigment, if it could be obtained from a natural ore, is a very desirable one, but the mines that furnished it in former days have long been exhausted as a commercial source of supply, and the copperas reds have taken its place. These contains a large amount of sulphuric acid loosely held in combination (see Analysis, Chapter III), and their use with red lead is as disastrons to the point as the direct effect of hydric sulphide.

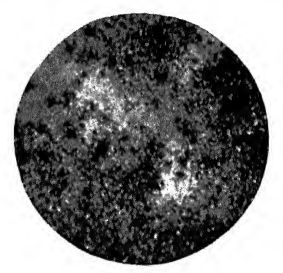
The effect of the sulphur element in both cases is greatly influenced by heat, a few weeks' exposure to hot summer temperatures being all that is necessary to destroy the conting by remiering it buildle and easily removed by the hand, or else causing it to perl in strips. This action of the sulphur is sometimes said not to be due to a change for the worse in the red-lead atoms, but to the change in the sehicle. But the effect of sulphur upon dried linear find in almost mil, and the oil-film is always nearly transparent of needs, whatever pigment may be associated with it. Any change in the payment is denoted by a change of color in the paint, whatever may be the cause of the change. The change in the pigment sample shows through the thin film of dried oil quite as reachiv as though it were of glass, and generally indicates a speedy dissolution of the conting. In many mills and workshops all of these conditions heat, sulphurous fumes, and saturated atmospheric elements ware present, and test lead coatings in such locations are short-lived.

The sulphur element, whether in the oil or the red hard, or from any other source, renders the paint liable to dry on the outface only, and the inner portion of the oil that encloses the pignish african remains soft and, therefore, more sensitive to my destructive influences that reach the coating.

From 28 to 30 pounds of red lead to a gallon of oil are mercuary to make a good red-lead paint, for even when well ground it is halder to streak, curdle, or run, and is difficult for the painters to operate The bulk of the red lead is so small compared with an equal weight of any other pigment per unit of covered surface, that the atomic of the red lead are well housed in the oil and letter presented. Hence, when the conditions are forwardle for

worked or brushed out in spreading. The porous character would disappear when spread on any surface other than glass.

The United States and other Governments have favored in the past a mixture of two or three parts of red lead and one of zinc oxide for the protective covering for lighthouses and seacoast iron structures. These coatings are harder than red lead alone, and better resist the action of salt-water spray, fog, and the abrasion from the sand-blast usual in such locations. But the result of some thirty



Parternierrograph . test of rest had and rel taken from a paint-put manel nergy no nel complete . I locate the profession of the profession of the contract of the trianguare of its operto, althought to the eye it leads outel. (M. Terli.)

years' experience with these maxtures has led to their abundonment for the reason that the coade of any in the centing changed to a carhonate of one, and he are merene of solume disrupted the dried conting, exposed the instruct, and the increase in the corresion was markedly greater than with red lead above, or red lead and silica. ear real least nearly extra large land love ter

Red-lead compositions are extensively advertised to keep indefinitely without setting, and that are ready for use at any time without further mixing or preparation. In all such mixtures, the red lead,



Fig. 11.—Photomicrograph×100 of a film of dried paint taken from an irrent pillar showing rust blisters. The dark spot is red lead and a incourer runs through the centre. The sine oxide and white lead are white still are intact. (M. Toch.)

or the oil, or both, are adulterated and will be found to be comparatively short-lived and unreliable whatever may be the guarantee, which in general lays more stress upon the extraordinary large nurface that can be covered than the permanent character of the coating. As well expect a hydraulic cement ready mixed to be a suitable article for engineering use as an "always-ready" red-lead paint.

The setting or solidification of a pure or nominally pure residend paint is a characteristic chemical union between the oil and the lead, and without this action the paint is worthless. This chemical action is sought to be simulated in all compound paints by the liberal tree of driers either incorporated in the vehicle by heat or by being introduced through the bung-hole of the barrel. The setting must take place eventually, and the better paint will be the one in which it is definitely provided for, and not left to the burdayard constitution.

One gill of crude mineral oil or heavy refined petroleum added to a gallon of red lead paint will delay the setting of it indefinitely. It will dry superficially, as the oxidizing power of the red lead will ensure that essential, but the petroleum will always remain viseld in the coating and eventually destroy it by peeling soon after an exposure to a strong similable or heat, following or followed by a lower damp temperature or a storm.

Red lead, either in the form of a pigment or paste, when quoted as being "second quality," can be regarded not only with suspicion, but with a certainty that it is greatly adulterated or poorly exidized from impure lead, or not properly washed or pulyerized.

First class manufacturers of red lead have no second-quality product that they are willing to have bear their brand or seal. There are a large number of red lead corrolers in the United States, and to the author's knowledge only one of the number advertises a second-class product. It may also be of interest to note that the United States Bureau of Construction, in its orders for red lead, specifies the make of one corroler as the standard of quality to which all tenders must conform.

A good red lead as it comes from the manufacturer is finely pulverized, as this point in a great measure governs the setting and running tercep, crawl, or curling the painters wall it). The atoms should be opaque, which indicates a good covering or light dispersing power. If the atoms are crystalline and more or less translinear, the paint will have a tendency to "tack." This effect does not always induste that the pigment is deficient in other respects to form a durable coating, for the "tack" is sometimes due to the quality of the oil, and that the red-lead manufacturer has seldom anything to do with

The adulteration of red lead and bitharge can be realily ascertained by digiting a sample in a warm adultion of nitrie acid; the adulterants will remain undesceled

Builing hydrochdoric acid will extract the iron oxide from the

Salt creates a chemical action on red lead that is hable to blister the coating and reduce the red lead to a metallic state.

Grinshaw recommends a mixture of red lead with painters' sizing to cover pine knots or yellow pine woodwork, instead of the usual shellae varnish. It forms a heavier coating than shellar, is equally or more resistant to the pitch, and is less liable to blacker.

equally or more resistant to the pitch, and is less fathe to brace.

A gallon of pure linseed-oil will require not less than 20 pounds as a minimum quantity of pure red lead to 30 pounds as a maximum quantity for a reliable red-lead paint which will cover trem 750 to 1200 square feet of metallic surface. These quantities of material at once remove red-lead paint from any companion of cost with the oxide-of-iron and many mixed paints, principally in the form of proprietary goods, the ingredients of which are only known to the makers, and the character and performance of which will start in quite as criatic a manner as the price paid for them.

The protective qualities of a well-oxidized pure red had and a

pure oil paint, properly applied to any structure under any expecture, except to the action of hydric-sulphide gas, cannot be care and. But what effect other than failure of it can be expected when a proventment engineer in charge of an important hydraula construction, after cleaning the metal part of the work by the sand libert, round it with the following paint? "Red lend, 10 pounds, torsed with their pints of water to one gallon of raw lineard oil for the first rout, and for the second conting, red lend, 10 pounds, three parts of water, three ounces of lampblack mixed with enough trapsation for each appearance of paste, and one gallon of raw lineard oil. It was found never any to first moisten the red lend with water to prevent the paint from streaking and sagging. Without the water, a large proportion of turpentine and drier would have been necessary, and this user considered injurious to the life of the paint. In warm weather a slightly less quantity of red lend could be used" for more water?

Many kindred examples of such "how not to protect" structures can be cited, none, however, more conspicuous than the above when the engineer in charge and the character of the work are considered.

The substitution of water for turpentine in the amount hate noted in order to molong the life of the point will be analysis to the solution of the solution of

financial part of their enterprise will seem in comparison but an insignificant pool. (The effects of watered oil is further considered in Chapter NAV.)

Lithurge, PhO (Protoxide of lead). Specific gravity, 8,50 to 9,00.

Lithurge as the first product in the oxidation of metallic lead to form red lead has been described. Another source of lithurge is from the scum of melted lead or that from the smelting of silver-bearing ores. It is formed as an oxide by exposing to the roasting heat of a furnace the slag, or "matte," that on cooling forms into white or take lithurge. The part that hardens hist is called "Massicot" or "levigated" lithurge, and is ground in water, dried, and made reads for the market. It is a yellowish red substance or an amorphous powder, and eristallizes in fine six sided scales or plates, it is a yellow or reddish protoxide of lead, partially fused and semi-transportent. The sellow is the fused or hard pieces that require to be ground and levigated; the red atoms are the flakes. The difference in the color areas from the mechanical condition resulting from the matnet and difference in exchanged condition resulting from the matnet and difference in exchanged product, a rapid cooling giving a yellowich redeet, a slow reading a reddich one;

By analysis it consists of the protoxide of had, (PhD),

the eight, burter mer , ... bie Ind , bur wie bert, od immenthe brief.

to the first of the transfer of the conditional areas, easily, exposer, autimizing bimuth, etc.

4 Mar. " flato " ersoners silvas, and christian sund. 43 Mar. " fraces " livia

The operial furnaces employed and the manipulations of the charge during the heating and cooling processes applicable to the manifacture of red lead are requisits for a reliable lithrage; also the name case in conduct and the subsequent operations to prepare it for a prement or for other uses. Its integrits when used in a paint is affected by the same carries that affect a red lead coating. It is adulterated if provide in a more barefaced manuer than any red lead

through maneral in analy from the lithrage "Manufact," also from white lead in most us the line

The carbonic acid in the white lead is expense, anyting a protocole of lead which absorbs more oxygen and produces a red lead of a lighter color than that made from litharge by tensors that the oxidation is more complete.

Paris red is prepared by roasting the earlierate of lead to a lithinge, the difference between the Orange mineral and Paris red pagaments being that the latter retains a little carbonic acid in its composition due to the different degree of heat employed in the furnace and the manner of cooling the product. Vermilieratte is an energy red pigment formed from the oxide of lead.

CHAPTER V.

WHITE BEAD.

White Lead, PhCO₈ (Hydrated Carbonate of Lead). Specific gravity, 6 465 to 6 5an,

The native anhydrous meta carbonate of lead, (PbCO_a), called white-lead ore or cerussite, when pure is found in cohorless crystals of the trimetric system. It is found in commercial quantities in all parts of the world where mineral lead ores are mined for smelting purposes. Plus mentions the use of a native ceruse found on the lands of Theodotus at Suvena.

The prote sulphale of lead, (PlS), is the blue-lead ore (Galein) and is the principal source for the supply of metallic lead. White lead and the red oxide of lead are next to the oxide of iron, order, umber, and siema, the object known pigments. Dioscorides (n.c. 400), Pliny, and Vetruxuis all mention the production of white lead by expessing metallic lead to the vapor of vinegar, giving the product the name of "Cerusa" and "Cerusa". Bergman in 1775 localized it as a carbonate of lead metend of an acctate, as it had before been considered.

White lead was used by the Egyptians as a cosmetic long before its employment for a juginest

The mining and meeting of lead ore to produce metallic lead were practiced by the Chinese 2000 years in . In the smelting of lead ore large quantities of the lead are expliced to the red lead "minimm," the use of which as a particult anticlates the knowledge of producing white lead to correspon. Moses commanded the Israelites to purify lead teatled optimists for

The returned amount of white lead is repulsived by the severalled

was almost synonymous with that of Helland, and explicitly was introduced by Hollanders, hence the name "Intal Process Lead." In this process thin perforated sheets of lead are expressed in gallipata

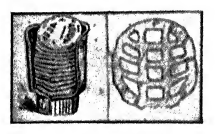


Fig. 12. Short-lead buckles and just

containing a weak solution of acetic acid (water with 23 parts of strong acid) or common elder vinegar. The puts are placed in long tiers. each tier being loosely covered with learning and starked in large tities. bers, 9000 to 10,000 pots containing 60 or more term of metallic lend, The bed of pots is then embedded in tandark, navidant, stable litter, etc., that ferments and som raises the temperature of the mace to 140° or 165° F. A quantity of vinegar containing det parameter of strong acid converts 2 to 21 tons of lead into the carbonate of lead to almost 100 days. The only attention the hed requires during the person of corrosion is to control the temperature of the mass to regulating the admission of the air to the interior of the back his representative or a housing the apertures left for that purpose. The controller in practically completed at the end of 60 days, but the least is ad light operation grassits. so it is the practice to allow the heds to remain universities for the or the days more, in which time the lend acquires a proper denser's lead is allowed to remain in the bests too long, and for the assenttion, it is liable to become crystalline and transparent and wall be of posses covering power. Care is necessary in the use of stable letter or that from flesh-eating animals, as they are liable to charge the white carbonate of lead as it forms into a dark sulphide of lead from the pulphurous hydrogen evolved in the decomposition of the manner

At the time of stacking the air in the twel contains almost the parts of oxygen; after 2 weeks it will and

through a series of rolls, beaters, and screens. The corroded lead is then mixed with water and ground between buhr-stones to an impulpable powder. Generally this part of the process is omitted by the quick-process lead manufacturers, because of the fine state of division to which it is necessary to reduce the metallic lead for these processes. The uncorroded particles are so intimately associated with the earbonate that they are indifferently eliminated in the separator, and if run through the water stones, will cover the face of the stones with a conting of metallic lead that soon impairs their grinding power and imparts a dark color to the product.

If the preliminary washing before grinding is not thoroughly done to free it from the acetic and (which is a drier) the powdered carbonate will dry in grains and hungs, and it may contain partly corroded or pure-lead particles, in which case the corrosion of them will proceed in the paint coating from the carbona and in the atmosphere. Added adulterants of any nature cannot prevent this secondary corrosion.

Silver in the metallic lead produces a pinkish east in the corroded lead; while bestingth inchines it to a dark or gray color. Antinony, arsenic, iron, zinc, and other metals also have a great effect on the color of the corroded lead.

After griding, the mixed carbonate and water is mechanically floated to remove any coarse particles, then jumped into large settling-tanks, where it is double washed with pure soft water and biearbonate of soda in solution to neutralize any trace of the acetic acid that may be present. After giving the lead time to settle in these tanks, the water is drawn off, and the judy lead, carrying about 24 per cent by weight of water, is jumped to large shallow copper drying-pans and the water exapitated. This drying process requires from 6 to 8 days, the temperature of the drying month being kept at from 140° to 160° F. The lead product when it leaves the drying-pans is judyerized and marketed as dry white lead or ground in buhr-stones with linsered oil for a paste or mixed paint.

A modification of the 19stch Process Lead known as "Pulp Procum Lead," consists of taking the pulp lead from the settling-tanks and placing it is a tank of linewish of and subjecting the mixture to a tion with the lead. Pulp lead is decided in advisor, which it is equently ground it does not bring it up to the folial review of a white-lead product. Pulp leads are inclined to if the interest from the same lead submitted to the full process of divide the rest of the interest ing. They are more uncertain in taking that was higher applied in frosty weather or on exposed situations, are provided to park. They are require more driers to aid in driving off the supplies matter. Then has price is all that gives them a market.

By the "Old Dutch Process" the lead is mather confirmed nor carbonated at the expense of the acturactal fibration the rived from the air, and the embonic and from the two buts, or other fermenting source. The vapor from the model closest waster a coaper orated by the heat of fermentation metels served to discolar the oxide of lead as it forms, converting it into a big is actuate which is again decomposed by the embonic and, the acceptable forms that, is, set free to act upon another portion of the lead

This is shown to be the mode of netters less a residence reason of corrosion, in which the protoxide of lead of last a residence is track with water containing about 1 per cent of the territal acceptance of lead (sugar of lead, PhO,C₄H_aC₄) and a current of carbonic acceptance are converted over it, the lithurge being quickly converted mater and excellent swhete lead.

There are many modifications of the "tild 1600". From There are referred to hereafter; all intended to improve the position of corrosion, and avoid the deleterious efficient fits when evolved from the corroding lead, upon the manufacture and consists and products by any process, from the lead one to a paper of a consist ingly detrimental to the health of all persons constructed activities the best-known precautions.

When honestly and thoroughly done through the hour of are of operations called the "Old Dutch Propose" the product a large smooth, and homogeneous in character as any known on the can be used with little or no waste. It is party desired to a sulphur compounds, which invariably change head there as an arrest to a sulphide, to the detriment of the color and life of the party.

the stack may differ from that at the side walls, where more moisture is present. An excess of most tire gives the grains a sugary appearance. The eveniness of temperature in the stack, due to many causes, also the time of the year that the corresion is effected, governs the quantity and quality of the product.

Commercial white lead is often inadequately corrolled, washed, and ground. Pieces of uncorroded lead, tan bark, and other substances from the ferment packing, are memperated in the pigment. subsceptent decomposition of which in the mixed paste or paint discolors it and shortens the life of the paint. To such an extent is the careless correspon of lead practiced, that the brain of "Ohl Initely Process" is called in decision "the happy godneky process" by the advantaged the rescalled "quark process". It is no longer a caternal of perfection in manufacture and parity, unless chiminal directly from reportable manufacturers or dealers. Nevertheless, white lead paste has held its former excellence and in some cases has been improved. It is also also to say that but few correlers continued to an the till Parch Process, however desirable it may In the loss of the report were. Los no doing residing to the confidence that implified and usees of paint, have no its ments, which have been

Some of the resilence process of corrector consist in the reduction of metallic lead action of areas or stand dark by which it is riven into small particles, greatly more and the authors exposed to the action of the action of the action of the action and there exposed to the action of the action and there exposed to the action of the action and the action action of the action o

The province by the claster has processed in not cold contract in composition but in some cases is decidedly resoluting mulcarries the water of crossallications that is afterwards set free in the grinling

oxide into a solution of neutral plumbic acetate (sugar of lead), is a white insoluble, anhydrous powder called "precipitated white lead." It is of comparatively recent use and is favorably reported upon for a pigment.

The hydrated carbonates of lead, formed by the direct action of carbonic acid on the hydrate of lead (Pb.(OH)₂), differs from the precipitated carbonates in being amorphous and perfectly opaque; whereas the precipitated carbonate is an aggregate of minute transparent crystalline grains. Hence the former are the best pigments; their greater opacity gives what the painters call "body."

In the German, Austrian, or Chamber processes of corresion, the

lead is used in sheets 1"×8"×12", 1800 or 2000 sheets in a box, 8 boxes to a chamber that may contain 12 to 24 tons of lead. The walls of the chamber are lined with metal and heated by steam. The carbonic-acid gas is made by the fermentation of vinegar, yeast, and other substances, ammonia, phosphate of magnesia, etc., being added to hasten the fermentation. Carbonic acid from burning charcoal and other methods are employed for generating the carbonic-acid gas in great volume, for a quick corroding vapor to fill the chamber.

The Kremser white or Klangenfurt, a German corrosion process,

uses the vinegar from dried grapes as an excitant to corrosion. The best quality of this process lead is claimed to be whiter than the "Old Dutch Process" leads and to cover equally as well.

Krem's or Crem's white is a poorer quality of the "Kremser process" lead.

Kremnitz white is a product from Kremnitz's (German) dry precipitation process.

Flake-white is a pure white lead in a scaly form rather than as crystals or grains—the usual form from the Dutch process. It lacks opacity or covering power.

The Clichy or French process is the principal quick-corresion process used in France. The product is known as Ceruse de Clechy. It is entirely different from the other decomposition or precipitation processes mentioned before. The white lead is formed by passing carbonic-acid gas for 12 to 14 hours through a sugar of head or list.

formed is more or less crystalline, loose or coarse in grain. It takes up less oil than the Old Dutch Process leads, allows more light to pass through it, hence does not cover nearly so well.

Greneberg's (German) process consists of the action of rarbonic acid on finely divided lead and litharge while being rolled constantly in tight metallic cylinders. The mechanical friction aids the corresion at the expense of the purity and durability of the product, though there is less exposure of the workman to the corrosion fumes in this part of the process.

Milner's (English) process produces white lead in two days by the action of carbonic neid on oxychloride of lead (litharge) by grinding them together with common salt in water.

Pattison's (English) lead is a wet precipitation product—the oxychloride of lead, made by the action of muriatic acid on galem (lead ore).

The Carter (American) process is a modification or an improvement on the Kremnitz (German) process. Metallic lead is unlted, and while molten is riven into fine particles, like flour, by a jet of highpressure superhented steam. This amorphous powder, of a steelgray or dark-blue color, is charged into a revolving cylinder 5 to 7 feet in diameter by 8 to 12 feet long. One end of the cylinder is conneeted to an exhaust-fan and the other end to a flue leading from a furnace where carbonic-neid gas is generated from burning charcoal. Generally the products of combustion from a coke fire under the steam-boiler of the plant are used for the corrolling gas, the furnace gases having been washed and parified to free them from any sulphur present. The temperature of the revolving cylinder and the charge of powdered lead is kept at about 140° F, during the process. Dilute acctic acid and hot water are sprayed into the chamber at different times during the corresion process, the stage of which is always accessible for inspection by removing samples of the lead without interrupting the chemical action of corresion. The agitation or turning-over of the lead and its exposure to the heat is constant during the process. About 95 per cent of the lead is changed to white lead by this process the care used to eliminate the acid the more reliable will be the product.

The products of all of the above processes, as well as of many other quick processes, vary in some degree of quality or form of the white-lead atoms from that of the "Old Dutch Process." The latter can be said to rank as the standard for purity, fineness, and all other qualities which are indifferently imitated in most of the quick-process products, the Carter being probably the best of them.

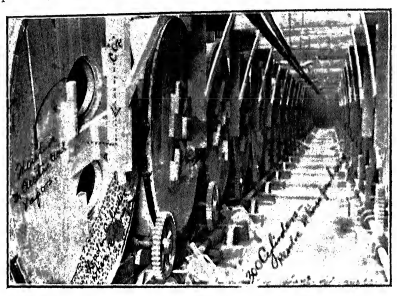


Fig. 13.—The Carter process for manufacturing white lead,

Many of the quick-process leads contain acrtic and carbonic acids; the former, being added in excess of the amount necessary for a natural rate of corrosion by the old methods, remains in the corroded product and requires a more thorough washing to remove it than it customarily receives. The acid element is often strong enough to redden litmus-paper, which would be discovered if the dry white-lead powder could be obtained to make the test. The acid causes loss of opacity and rapid chalking. The corrosion of the lead is simply removed from the corroding stack to the traint contains.

The report of the Committee of Experts appointed by the English

of the coating. Acetic acid and uncorroded lead, left by imperfect washing and grinding, are frequently present in commercial white lends. Ten per cent of lend acetate is often found in the "flakewhites," Most of the quick-process or impure leads come to the market in some form of "whites" with a misleading trade-mark.

Home Secretary to investigate the manufacture of lead products, as to the character and quality of the product, also their effect upon the health of the workmen, was: That they visited forty-six establishments using various processes for manufacturing white-lead pigments, all of which were dangerous to the health of persons so employed, and while some of the substitute leads were cheaper to make, and possibly a little less injurious to the workmen, their products were far from equalling in quality those from the "Old Dutch Process," and they could not recommend either the processes or products as against the "Old Dutch Process" lends, Baryta white is prepared from the native sulplante of barium, or

from the earbonate of baryta, artificially treated with sulphuric acid. (See Barytes, Inert Pigments, Chapter NVIII.)

Krem's, Nottingham, and Newcastle whites are pure white leads differing only in the process by which they are made. Humburg, Holland, and many foreign-made whites contain from 3 to 60 per cent of barytes and chalk, and are adulterated compounds of white lend. Verice white generally consists of equal parts of white lead and barvies. All pastes and mixed paints classified and marketed as "whites" are usually only adulterations of white lead, and no responsible and honest corroders of white lead ever so denominate their products. The name "white," whatever its trade prefix, should usually be viewed with suspicion of its quality. (See tests of white lend on the following pages.)

About 110,000 tons of metallic lead are annually corroded to white lead, in the United States, by the various processes, or about ope-third of the total production of the metal product.

There are twenty-two manufacturers using the "Old Dutch Process" in the United States, and five using the "Quick Process." The first white lead corroded in the United States was by Samuel afterward.

Electrolytic White Lead.*

This process is a radical departure from all of the other processes for producing white lead, in not employing acetic acid, but by acting upon the lead in the form of pigs with nitric acid, which is generated by electricity. The process consists of four consecutive steps:

First. The electrical preparation of nitric acid and sedium by-

droxide.

Second. The action of the nitric acid on the metallic lead forming lead nitrate, $Pb(NO_2)_2 + H_2$.

Third. The reaction of lead nitrate and sodium hydrate to form

lead hydroxide, viz.: $Pb(NO_2)_2 + 2NaOH - Pb(OH)_2 + 2NaNO_2$.

Fourth. The combination of the lead hydroxide and softum bienrabonate to form lead carbonate, Pb(OH)₂ + HNaCO₂ + NaOH + H₂O₂

Reactions 2 and 3 may not take place strictly as given, which are the theoretical combinations, but some approximate reactions are had, for the extra hydrogen present is liberated at the electrode.

The chemical operations in the process are briefly:

First, a solution of nitrate of sodium (NaNt) is decomposed by an electric current from a dynamo, the strength of the solution not being important—10° Baumé or one pound per gallon suffices. This solution is put in a series of wooden cells divided into two compartments by porous partitions. At the positive electrode is fastened a pig of lead, at the negative a sheet of copper. On applying the current the nitrate of sodium is decomposed according to the equation 1 given, nitric acid collecting at the positive electrode and sodium hydroxide at the negative. The nitric acid at once attacks the lead and forms lead nitrate, which dissolves (equation 2) the hydrate of sodium, producing no effect upon the copper at the negative pole.

Finally, both solutions are separately drawn off and mixed as desired, in quantitative proportions in any suitable vessel. The result, as shown in equation 3, gives the lead hydroxide as a white amorphous precipitate and leaves the nitrate of solution in addition.

small, and a little additional fresh sodium hydrate restores its strength. The lead-hydrate precipitate (Pb(OID₂) is then filtered from the sodium hydrate by a rotary separator, and the nitrate of sodium returned to the original reservoir.

The fourth step is in some respects the most interesting of all, and consists in adding to the lead hydroxide a solution of bicarbonate of soda (or the normal carbonate). Reaction 4 at once takes place. It will be noticed that the sodium hydroxide is the product in solution, and lead carbonate the precipitate.

The sodium hydroxide removes most of the impurities, if there are any, in the hydrate of lead. It dissolves any salts of alumina or zinc present, and it removes the organic matter. These impurities appear in the solution, leaving the precipitated lead remarkably fine and white. The hydroxide of sodium is again converted into bicarbonate by passing carbonic acid into it, and this is used again. Thus the main agents in each of the two principal steps, the nitrate of sodium and the bicarbonate of sodium, are made to do duty over and over again with but slight additions to restore the strength.

The use of free nitric acid in the process is objectionable, as under the influence of electricity it breaks up with intolerable funnes; also for other reasons. Acetic acid is also objectionable for the same reasons, hence the recourse to sodium or potassium nitrates for the reactions.

The cost of white lead by this process is but a fraction of that by the "Old Dutch Process," as the lead is used as it comes from the smelting furnace in pigs and requires no remelting or casting into buckles or shreds, as in the corrosion processes, and the whole process is complete in a day, or, for that matter, in an hour, as all of the reactions take place rapidly, if not instantaneously, no free acids are used, and the sodium compounds are recovered, as noted.

The texture of the lead product is almost molecular in fineness and does not require grinding, it being so fine that it remains suspended in the water for a long time, and in order to filter it a special brand of cloth is used, as filter-paper would scarcely retain it.

Its covering power applied side by side with the Dutch Process lead appears to be equal to it, possibly a little better, but never found. lead to any great extent remains for time to determine. The French or Chehy process lead, or "Chehy white," was thought at first to be a revolutionary one, but the product finally process to be decidedly inferior to the Unitch Process lead, from its crystalline character. It does not give the opicity or body, or operad as well inder the brigh, or cover as much surface as the Dirich Process lead.

The "Intch Process" lead forms a globular atom, we two atoms of the carbonate, Phitt, and one atom of the hydrate of lead, Phit III), but this composition does not always appear to be of ear stant quality, as much depends upon the care given during the correspondent of the process.

Lungert tan etrieben, aus fiber tan altieben und biter und beiter in ballabler jedeneungebereben mester geft annergen. There area all beer nen and beer nan bet beer ernelle erne geleeften bei eine eine eine beiden ber bergegentele. anne ragmete ther begerthreite eif ifin geneigendenteit. Does entfinan egrechtebein gaf thermer fober fergeben ibre agreeber bonebeller, indant biben erdgeftenben na fod. Eine nemer Bedrungenger in the bereit education about Bruch ber ber bereit bereit bereit bereit bereit bereit ben ben ben The atomic of the one form are part to be from , to be not be an interest to be सहित्यक्षक्रक में ताह , प्रकार है कहा में है जबर मुझक का है के प्रकार के में के किए का का का का का का का का मा requestinger calmouther maintains. Its ther " thestails I to me more " levisite in their satisfies of me of merely ungerundent, blur erbeiter gebeiter gebeiterlich, erbeit ber blieb bie ebreit ber geneubern bneite manif peranguangungang big after demonat visura agres garang manib ving agreeing diang bearing [kergkalin - [[]] ker ein noth toll tallanner foren eil retror harret Labernerth Floren thin Hindre (Pour tolland nother teather eighte son talland eile (und night, bonn begindlichen Grenne Bobin for it benieb finer genriebent inn liften mebnichen nich beimer grand magnatin, grandlung ind tarette, jam Grennbauel eine ernenfound aus biber genouelanauer jane auge genrgin bentere langel eine bered leine und fleer untdeteter begebet elemmermetetet ein wurden der bereit eine Marganian Abanan genenen unen unbabagt ignacabatin, janait fonnte ibe fannt bereget ber auf ban ann an Abber Serfagetter ibn bereit Aber's mie un not.

If he the electrolate present at a proposible can it is classically to produce a pitte carbonate of lead, or a mice time of the carbonate and hadrate of lead in an proposition required, and the product process to be fine and plobular motival of coarse graniclar or erastalline, there should be not disable regarding to morate, but the few hundred tone than far spread do not affect sufficient data for a wholesale abandonment of the 19th I butch Process" with its venturies of established requiration.

or roofs. The fine powders fall down and are collected for pigments, and require no washing. The uncondensed vapors escape through the cloth screens to the atmosphere or stack. A saving of 50 per cent is claimed for the process over that of the "Old Dutch Process," but like all other quick-process leads, the merits of the product are yet to be established.

There never has been any difficulty in quickly corroding or oxidizing metallic lead or zine for use as a pigment. The difficulty with all quick-process leads is in the quality of the product, and, though the processes are patented from Dan to Beersheba, the ormate devices and claims of the patent-office document do not put the wearing and other desirable qualities into the product that the patent claims allege to be there.

There is nothing electrolytic in the Bailey process. The melting of the lead by an electric current previous to or during the action of the corroding gases produces no electrolysis in the molten metal, the heating of which could be as readily and more cheaply done by a coal or other fire. The collection of the metallic vapors by screens is an old method of pigment manufacture—in use for the production of sublimated lead, as described in Chapter VI, and is void of electrolytic action.

CHAPTER VI.

WHAT CONSTITUTES A GOOD WHITE DEAD.

MULDER's and Phillips's analyses determined that there are three varieties of white lead in the best classes of that product experimented with under the general formula of the hydrated carbonate, Phi's, viz.;

A properly corroded white lead should contain exide of lead combined with water ewater lead (hydrate of lead, Ph(OH), i, from 25 to 32 per cent, which in its effect upon the vehicle is similar to red lead; it absorbs exygen and hardens the coating by converting a part of the oil into a soap that has no covering power whatever. The other 75 to 68 per cent should be exide of lead combined with carbonic acid (carbonic-acid lead), that really injures the oil in the paint, but gives all of the covering properties that the paint passenes. The chalking of white-lead paint is due to this 75 per cent of carbonic acid lead. A paint composed wholly of carbonic-acid lead will, in a short time, chalk as freely as a whitewash. The carbonic-acid lead gives the whiteness or color, the water lead, the hardness or durability to the coating.

In the following table from Heiss's experiments,

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In Prof. Hurst's analyses of four samples of "Old Dutch Process" white lead, the carbonate ranged from 65.35 to 72.45 per cent, the hydrate from 25.19 to 36.14 per cent, and the moisture 0.42 per cent to nil.

Mr. Converse's analyses of five samples of the best brands of American "Old Dutch Process" leads gave

er be		100100000000000000000000000000000000000	11 - 11 - 24 AND	ominge y formulae confequences and addresses	TO STEED THE LOSSING CONCENSION AND ADDRESS OF THE PARTY	*
" hydrati " oxide	ate		79,37 19,80 0,21	78,58 20,11 1,48	77.98 20,60 1,48	69,96 30,19 0,07

A very hard white lead that contains no water lead will not harden when spread, but brush off like a lime whitewash. When this occurs where the necessary amount of oil has been used, the painter can be quite sure that the lead is a poorly corroded or quick-process lead, or a synthetically formed lead from an acid-solution process. Carbonate of lime (chalk, whiting), gypsum, and other so-called inert substances added to such a lead do not correct this chalking; they only disguise it for the time being, and increase the tendency of such lead paints to turn yellow and lessen their covering power.

All white lead on external exposures is liable to chalk, because it contains too much earbonic-acid lead as it comes from the manufacturer, or has taken another portion of carbonic acid from the atmosphere at the expense of the water lead, and formed a subcarbonate of lead (the chalk product, PbCO₂), or has too little oil in it when spread, or has not been well ground in the oil. Paints thus affected, if brushed over with a conting of white lead ground with an excess of oil, will prove to be more durable and less affected by future chalking than the original coating, or a new heavy coat from the same lead.

The formation of a lead soap in the ordinary process of grinding and mixing white-lead pastes or paints is a disputed point by paint chemists. But the lead hydroxide and the free linoleic acid in linsced-oil, if acetic acid is present in the white lead, will combine and form a lead-soap mixture. The paint containing this soap, on exposure to the weather, soon loses its lustre and will crumble or chalk.

The presence of lead some in many, if not the most, of white-lead

fuse character of the paint. With zine white of baryta write in on no such admixture is possible, for in these paints the oil will not saponify owing to the absence of an acid."

Mulder's and other experiments proved that the chalking of white lead was due solely to the absence of water lead in the pigment, unless the lead was badly adulterated, in which case this effect was directly

traceable to the adulterant used.

In general, all the latter-day-process white leads are more inclined to chalk than the "Dutch Process" leads. This arises (as stated before) from the smaller amount of water lead in their composition, and being of a crystalline instead of a globular form. In the granding process these crystals or grains are broken down, and the combined or formative water necessary for their existence in the form of crystals is dispersed, rendering the broken lead atoms more sensitive to the attack of the atmospheric carbonic-neid element, that finds in their sharp angular form a more favorable surface to act upon. Unity a comparatively slight action of the carbonic acid on the freshly crushed atoms of the lead is required to change it to the subcarbonate, and leave it free to be brushed off by friction or washed out by storms.

Old white lead, or that which has been ground in oil to a paste for

Old white lead, or that which has been ground in oil to a paste for a year or more, chalks decidedly less than recently corrolled lead, by whatever process it is made. The atoms of the pigment in the case of old lead have had time to release themselves from the tension due to their formation and grinding. Adulterations do not prevent these inexorable chemical changes in the lead pigment; they only increase the disintegrating action.

If oxide of zine is used to give hardness in place of the water lead in paints exposed in open air, the atmospheric moisture and carbonic acid changes the oxide of zine (ZnO) to a zine carbonate (ZnO), whose volume is nearly double that of the oxide from which it was formed in the hardened paint, and peeling takes place instead of chalking. The cheaper forms of zine oxide contain zine sulphite, which blackens the paint and otherwise injures the conting

If gypsum, barytes, or silien are used for the adulteration, they lack both in covering and light-dispersing (coloring) power, and from their sharp, angular, or irregular form as pigment-atoms, do not bond themselves in the cit of a silicit or a silici

if they do not actually increase its tendency towards that change, as the sharp, rough surface of the paint containing these adulterants holds the atmospheric moisture and gases closer and longer for their action. (See Decay of Paint and Inert Pigments, Chapter XXVII.)

White lend unites thoroughly with the oil. Zinc white combines, but very slowly; barytes does not combine at all. All pigments that contain crystals or are granular are deficient in the light-dispersing power, even if they have the spreading quality. The granular character of quick-process white lend is its great weakness.

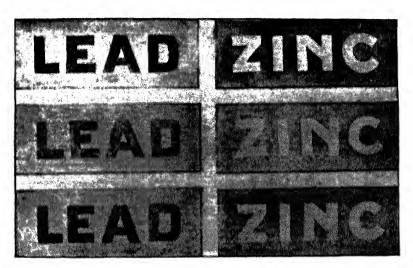


Fig. 14 Effects of sulphurous gases on white lend, Background in suc white.

Background in white lead.

Lettering in white lead.

The quality of the vehicle has much to do with chalking and the decay of white lead. A good linseed-oil will better preserve a poor, or an adulterated white lead, than a poor oil could the best of white lead.

Pure white lead is soluble in dilute intric acid. A sample treated with this reagent should pass entirely into solution, leaving no residue.

in the volume of the lead-atoms at each step in their progress from a carbonate to a sulphate that ensures a forced disintegration of the coatings. (See Decay of Paint, Chapter NXVII.)

The cause of white-lead paint, when sprend over darker colors, deteriorating after a short period and showing dark, is not alone owing to the changes mentioned above, but also that the fatty acids in the oil gradually expel the curbonic acid in the lead atoms and form a clear lead soap, through which the darker colors beneath show.

Adulteration of White Lead.

The adulteration of white lead under the forms of a paste or mixed paint has reached a point where the multitudinous trademarks under which they are marketed are positively of no value to determine their quality; the only safety lies in purchasing from responsible business firms of national reputation for the standard quality of their products and business methods.

A late examination of commercial white-lead pastes purchased in the open market resulted as follows:

Seventy-five different white-lead pastes, under 29 different trademarks and symbols, embellished with 14 qualifying adjectives, and made by 17 different manufacturers, plus I nuknown firm that furnished 14 different brands, were analyzed by 16 different analysts. The condensed results are, viz.:

Sixteen had no white lead in their composition, but were mixtures of barytes, silica, gypsum, zinc oxide, or whiting in some proportion of three or more of these substances.

Fifty-nine of the samples had white lead from 1.24 per cent to 47.62 per cent, and averaged for all 23.35 per cent of lead.

Seventy of the samples had barytes from 15.60 per cent to 86.37 per cent, and averaged for all 49.61 per cent. One or more of the above adulterants was used with white lead of uncertain quality for the balance of the paste.

In 5 of the samples silica replaced the barytes, otherwise the usual group was unbroken.

Seventy-five of the samples had evide of vine from same

pure white-lead product.

Thirty-two of the samples had sulphate of zinc in the paste from

84.85 to 0.88 per cent, averaging for all 22.09 per cent.

Five samples had chalk from 24.30 to 0.85 per cent, averaging for all 9.60 per cent.

Wherever carbonate or sulphate of lime was used it was at the expense of the white lead.

The adulteration of all of the brands, other than the 16 that had no white lead, ranged from 99 to 44 per cent, and averaged for the 59 pastes 80.4 per cent.

In 5 of the samples oxide of zine was mentioned as constituting a small part of the paste, but nothing was said about the 30 to 40 per cent of barytes in their composition, or of a like amount of the other adulterants, or the small amount of white lead.

One sample had a notice that \$1000 would be paid if the white lead in it was not pure. It had no white lead in it.

One sample offered \$100, same conditions as above. It was of the same character. No white lead in it.

One sample had a \$250 penalty for any inditation of the lead in it, or of the label on the package. It had 99 per cent of adulterants. Possibly a piece of the label might have fallen into the paste before analysis of it.

Probably the above adulteration record represents the larger number of commercial white lead and tinted-color paints in the market. Unfortunately, most of them get juggled off, and generally at the price of really first class paints; but so many people are willing to be humbugged to save a few cents on a pot of paint that it may be a mistake to subghten them on the subject. What the character and quality of the oils are that form an essential part of all these adulterated and misleadingly named compounds, one can readily imagine. Surely they are equally deceptive and uncliable.

Maize oil is used frequently to the amount of 25 per cent in the grinding of paint pastes and paints to prevent them from settling in the package. It is a non-siccative oil, and while not exerting any materially had influence in a paint, its use with linsced-oil in any quantity can only be considered as an adulterant. Its cost is about barytes, but zinc oxide covers beller; barytes gives weight that the zinc is deficient in. Some master painters advocate the use of barytes for the reason that it brightens dark colors and saves oil; but ignore the fact of its darkening light colors, also its tendency to yellow them from the sulphur element in its composition and its general deficiency in covering power. The advantages in any case do not warrant the use of from 50 to 80 per cent of this or any other substance which does not unite with the oil, and of themselves are unfit for a paint. As a rule, no responsible paint firms will offer such paints under their own names.

Zinc oxide is often mixed with white lend for other than adulteration purposes. It is added to correct the tendency of pure white lead to turn yellow from the action of sulphurous funces in the atmosphere. It makes the paint harder, and possibly prevents "chalking" to some extent, as there is not so much lead to chalk, but adds a tendency to peel on outside exposures—a result worse than chalking. It is difficult to get a homogeneous blending of the two pigments—or individually unlike in character, owing to their different processes of formation. Their mechanical association is far unlike that of a neutral whole, even with the best of supervising care in blending them.

Many of the difficulties in mixtures of white lead and zine oxide are overcome by the use of a natural admixture of the lead and zine in the form of a pigment known as sublined lead or white paint, hereafter described.

White Lead vs. Zine Paints.

Mr. G. R. Henderson, of the Norfolk and Western Railroad, reports a series of exposure tests to determine the efficiency of lead and zine paints. For the different uniterials he reaches the following results:

"Tin. The best results were obtained with the first cont of white lead and second coat of white zinc. The second coating of zinc gave generally the best results, and the second coating of lead the worst.

"Galvanized Iron. The same remarks apply to galvanized iron that are given for tin.

"Sheet Iron. The mixture of one-third white zine and two-thirds

matter whether the princing coats were white lead or white zine or mixed lead and zine. The lead second coating showed up the worst on this material, but in each case where the second coat was of zine, totally or partially, the paint was in a perfect condition.

- ⁹ White Pine. The remarks made relating to poplar apply to white pine also.
- "Yellow Pine. This material seems to be difficult to properly treat with paints; the best results were obtained with the first coat of lead, and the second coat of lead and zine mixed. Where the first coat was of lead and zine mixed, or entirely of zine, the results were poor throughout, which seems to indicate that as a general thing the lead is better for prinning out this material.
- "Conclusion. The lead priming and zine coating is generally good for tin, galvanized iron, poplar, and white pine. Sheet iron showed up best with both coats of mixed paints. Yellow pine appeared best with the first coat of lead and the second coat of lead and zine mixed. Comparing the materials which were painted, we find that generally poplar retains the paint better than white pine, and would, therefore, he preferred for siding on buildings, etc. Yellow pine seems to be the worst of all for this purpose. Black iron, as a whole, seems to retain the paint better than either tin or galvanized iron."

Tests for White Land.

There are many tests for the purity of white lead, a simple one being to put a globule of the paste or dry powder in a cavity formed in a piece of charcoal, and expane it to the heat of a common blowpipe, readily extemporated in any shop. If even 10 per cent of adulterants are present it will not be possible to melt the mass.

White lead is tested for barytes by dilute nitric acid, in which barytes is insoluble, while the white lead passes completely into the solution. Whiting and chalk are detected by the nitrate solution yielding a white precipitate with evalue or sulphuric acid or exalate of animonia, after having been treated with sulphuretted hydrogen or a hydrosulphuret to throw down the lead.

White lead dissolves with effervenence in hot hydrochloric acid as

	Conduct towards						
Substance.	Muriatic Acid.	Caustic Soda.	Heated before the Blowpipe.				
Whiting or chalk in any form.	Soluble with effervescence.	Unchanged.	Becomes incan- descent and turns tumeric brown after cooling.				
Commercial white leads. Pearl and other whites. Carbonate of lead, etc., quick-process leads.	Soluble with effervescence and deposition of small crystals.	Soluble with- out residue, or if of poor quality, 80 per cent will only be dissolved.	Coating formed on the charcoal is citron-yellow while hot, sulphur- yellow when cold; is easily fused with metallic beads.				
Pattison's white lead. Lead oxychloride.	Same as above.	Same as above.	Same as above.				
Zinc white or oxide.	Soluble, no ef- fervescence.	Soluble with- out residue.	Yellow while hot, white when cold.				
Antimony white. Antimonious-acid pigments.	Same as above.	Same as above.	White, easily volatilized, metallic globules which give off white smoke.				
Bone-ash, Bone- black. Phosphate of calcium, Ca ₄ (PO ₄) ₂ . (Basic steel furnace slag.)	Soluble after heating. Effer- vescent at first.	Unchanged.	Unchanged, but becomes incandes- cent.				
Barytes, Blanc- Fixe, mineral white, and other pigments of the native sul- phate of barium.	Unchanged.	Unchanged.	After ignition, if moistened with muriatic acid, gives off sulphuretted hydrogen vapor.				
Gypsum (Hydrated sulphate of calcium).	Unchanged.	Unchanged.	Incandes cent, like barytes. If heated in a tube, gives water vapors, etc.				
China, pipe, pot- ters', and other	Unchanged.	Unchanged.	Incandes cent, same as barytes				

lead, and the mass is allowed to stand a few hours, the temperature of the mass becomes so high that the oil is carbonized and colors the mass dark or even black. All white-lead paste is liable to turn brown if exposed freely to the air, hence should be kept closely covered, or water be kept on the paste when the package is opened.

Gmelin states "That the 'Old Dutch Process' white lead diffused through water, under the microscope appears as non-crystalline, round, and oval globules, .0001 and rarely .0003 or .0004 of an inch in diameter, while in the quick-process leads the globules are distinctively larger and more transparent and crystalline."

Dry white lead is tested by heating 100 grains red hot and stirring it for a moment. Its loss in weight by driving off the carbonic acid should be from 13 to 16 grains. If more or less loss is incurred, the lead is probably adulterated, and should be submitted to further test to determine the character of the adulterant, as shown in the table opposite.

Sublimed Lead, PbSO₄. Specific Gravity, 6.258.

Sublimed lead is a by-product obtained in the smelting of non-argentiferous lead ores. It is known in the trade as Joplin lead, from its place of manufacture, Joplin, Mo.; also as Picher lead, from the name of the manufacturing company. It is made in two colors—white, suitable for all purposes that the hydrated carbonate of lead is used for; and blue, which is a preferable color when used as a paint for iron. Both colors are used in the manufacture of india-rubber articles. The chemical composition of sublimed lead is sulphate and anhydrous oxide of lead, both amorphous; the good qualities of the white are also present in the blue-colored product. There is a small percentage of zinc in the Missouri lead ores, which in the process of smelting is converted into zinc oxide and is found in the sublimed-lead product, as may be seen in the analyses of a sample of the white product given on page 84.

The blue pigment owes its color to the lead sulphide and carbonaceous matter from the bituminous coal used as a fuel in the smelting-furnace.

Sublimed lead is prepared in special furnaces, the process being patented and known as the "Lewis and Bartlett Bag Process for

metallic constituents of the galena ore and "fume." The latter is drawn off by an exhaust-fan through a settling-chamber to a bag-house which contains a large number of woollen bags for filtering the fume out of the combustion gases. This "fume" is a lead-colored, impulpable powder known as "blue powder." It is ignited and allowed to burn for several hours, which converts it into white coherent crusts. These crusts, with some oxidized ores, settling in the flues and hearth slags, are next charged into a special furnace and exposed to a very hot coke fire. The products of this smelting are pig lead, slags poor in lead as waste material, and the "fume," which is now a perfectly white impulpable pigment suspended in the air. This is drawn through a series of cooling flues, where a further purification takes place, a part of the product settling and the action of the heat. The sublimed lead is now arrested by forcing the gases and lead into strainers of fine textile fabric, where the gases The process assures a more intinate combination of the vaporized

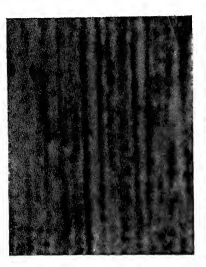
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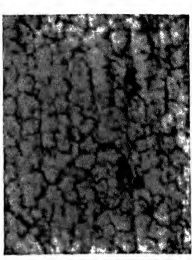
carrying down small quantities of any impurities that have escaped escape by filtration. The sublimed lead is taken from the strainers and is ready for the market. atoms of the lead and zine to form a neutral whole in which every atom is approximately of the same physical character and equally affected by any of the causes that injure a paint, than is possible in a pigment formed by a chain of partly chemical and partly mechanical operations acting upon a number of separate substances at different stages of the manufacturing process. Sublimed lead is absolutely free from soluble acids or sulphur, is amorphous, non-crystalline, and fine and smooth in the character of its atoms. Being a pyrogenicformed substance, it is not affected by heat or deleterious gases of

the atmosphere or manufactories; does not turn gray on long exposure to the simlight, and is not liable to chalk like correlled white lead. Though it dries firm and of almost ivory hardness it does not blister. crack, and peel like the oxide of zinc or mixtures of zinc oxide and corroded white lead. It is clastic, mixes thoroughly with the oil, and dries well without an excessive use of driers, either metallic or liquid. Pound for pound it covers better than white lead, and keeps its color

the extra contract of the cont

paint shows decay, the coating can be repainted without removing the old paint. Either the white or blue products of sublined lead will take a tint more uniformly than is possible with any mixtures of white lead and zine oxide incorporated by the usual grinding processes.





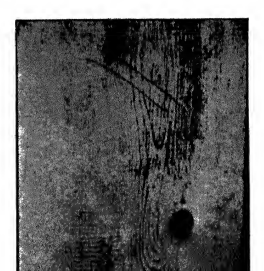
Frg. 15.—Sublimed lead.

Fig. 16. - Pure correled white lead.

The cuts are reproductions from a photograph of a picket fence, each alternate picket being painted with the different leads at the same time and by the same painter, using the same off as a thinner for the pastes, and the same driers, a separate brush being used with each paint. Three coats of each paint were applied, each being allowed to dry thoroughly hard before the next coating was spread. After an exposure of three years and one month the photographs were taken. The cuts are magnified ten times. The chemical reaction between the corroded white lead and the oil forming a lead or paint soap is clearly apparent. The shrinkage of the paint map has caused the coating to crack. Moisture has entered and language the bond of the paint to the covered surface, while the soluble character of the lead soap has caused the whole coating to "crace" and



Fig. 17. Sublimed lead.



streaking being due to the running down of the road dust when wetted by storms, no crazing being noticeable.

Fig. 17, sublimed lead, and Fig. 18, pure corroded white lead, are two-coat applications of the respective paints on a well-sensoned board after an exposure to the atmosphere for three years.

CHAPTER VII.

ZING OXIDE (ZnO) AND OTHER ZING PAINTS.

Metallic zinc (Zn), specific gravity, 6.86 cast; 7.14 to 7.20 rolled = 437.5 pounds per cubic foot; combining weight, 65.4; tensile strength, 5000 to 6000 pounds per square inch; electrical conductivity, 29.—It melts at 780° F, and begins to volutilize in the open air at 800° to 825° F.

Zine is electro-positive to copper and iron, whether in solution or in contact. In contact with iron or steel it forms a galvanic pile and decomposes with the evolution of hydrogen. It is used in this form to protect steam-boilers from corrosion. Each pound of zine decomposed evolves 5,6948 cubic inches of hydrogen that weigh 210,29 grains and develop an electrical or decomposing energy of 1,172 horse-power when used in a sulphuric-acid battery or 1,06 horse-power in a Bunsen or Grove battery.

This electrical energy, while not so strong as in the oxide or salts of zine in a battery form, is still present, and is recognized by painters as "movement in the paint," which is very marked with zine pigments, as will be hereafter explained.

Mallett's experiments determined that copper and zine plates in contact with iron increased the corresion of the iron 60 per cent. Copper alone in contact with iron, 40 per cent. Elsner found that the oxides of tin, zine, and iron used together in a paint set up galvanic action enough to crystallize the tin into flakes, which could then be rubbed off.

Zinc is associated with nearly all of the other metals as a mineral ore. It is roasted in special furnaces and by processes similar to those used in the reduction of lead ores for the metal or its oxides.

associated with it.

Zinc forms but one oxide, ZnO, composed of zinc 80.344 per cent and oxygen 19.656 per cent; specific gravity, 5.42.

Zinc oxide is produced by two methods: First, by the oxidation of the metal (French process), which gives a more dense and harder pigment than that prepared by the second method—the sublimation of zinc ore (American process). The product from the first process is preferable for straight oxide-of-zinc paint; the second is better for use in combination with other pigments. Both varieties grind and mix better with poppy-seed oil than with any other vehicle, and man ganese-borate drier should be used in preference to lead driers, which are liable to blacken the paint on exposure.

In both processes the zine is reduced to a vapor by the furnace heat (about 850° F.), and is exposed to a current of air that changes it to a flake, filament, or needle form, according to the care exercised in the process. Some of the particles of the metallic zine are upt to be carried over in the vapor unchanged, mixed with the atoms of earbon from the reduction fire. These carbon-atoms tend to give a gray color to the product, while the metallic-zine particles are subject to an attack from the carbonic acid in the atmosphere, forming the zine carbonate (ZnCO₂); this latter change occurring in the pigment after it is mixed into a paste or paint.

The French brand of zine oxide (La Vielle Montague Co.) is the best in the world, probably, due to the purity of the metallic zine, the care exercised in its reduction, and by the use of poppy-seed oil, with which the oxide is ground immediately after its formation.

Most of the American zine ore contains lead, tin, antimony, his muth, silver, etc., the oxidation of which in the reduction furnace produces white pigments, but they are all blackened by sulphaneous hydrogen, which affects the quality of the zine product, and aids in setting up the electrolytic action that all zine substances are sensitive to, as before stated. Sulphur is also present in many zine ores, and causes a yellow color in the oxide.

History of Zinc Oxide.

In 1701 a Propose about the second 111

It and not come into general use against write lead.

In 1844 Leclair, a Frenchman, made zine oxide and founded the Le Vielle Montague Zine Co. Leclair died in 1872. He received a gold medal from the Society for the Encouragement of the Arts, and was decorated with the Grand Cross of the Legion of Honor for having improved the practice of painters.

Leclair disquised the fact of the use of zine oxide as a pigment. His product was always sold as white lead, a precedent that mixed-paint immufacturers of the present day follow but too well if the analyses of commercial white leads given in Chapter VI are any indication.

The Leelair process consists of volatilizing the metallic zine in a retort, the vapors as they issue from it being met and mingled with a current of hot air from a blower which completely oxidizes them. The resulting products of combustion are led through a series of flues and chambers, where the zine oxide is deposited in the form of a flocculent, impulpable white powder ready for use as a pigment.

The American process for the sublimation of zincite ore into zinc oxide was invented by Mr. Samuel T. Jones in 1850, who constructed a furnace for effecting this purpose. This process was improved by Col. Samuel Wetherell for the purpose of working the franklinite ore from the New Jersey deposits.

Wetherell's invention embraced a special furnace and a process. The zinc ores are mixed with pulverized authracite coal, and charged into a closed furnace having a perforated grate, through which an air-blast furnishes the air necessary for the combustion of the coal and oxidation of the zinc. The vapors from the furnace are led through a number of flues and chambers, where the coarser particles are deposited, while the fine air-floated atoms of the zinc oxide pass on and are collected in a number of fine muslin bags, through which the combustion gases filter away to the atmosphere; the latter part of the process being similar to the Lewis and Bartlett bag process, used in the production of sublimed lead, described in Chapter VI. The franklinite and zincite ore found near Mt. Sterling, N. J., now furnish most of the American zinc oxide.

About 44,000 tons were produced in the United States in 1900,

color and quality of zinc oxide. In France its use has almost super seded white lead for interior house-painting, the Government prohibiting the use of white lead for this purpose.

Zinc oxide made from the ore is used more extensively than that made from the metal. The latter not only dries harder and is more brittle, but on large surfaces the difference in the whiteness of the coatings is very apparent and in favor of the mineral oxide. Zinc oxide mixed with water tends to collect in humps or masses. It should be thoroughly dry before being ground with the oil. It does not unite so thoroughly with oil as lead or iron pigments, nor dry as quickly.

One-third of one per cent of lithurge, added to the linseed oil in which zine oxide is ground, renders the paint more elastic and less liable to peel.

Mixtures of zine oxide, white lead, and ground silex barytes in the proportions of about one-third each, prove very durable in southern climates and seaconst exposures. The silex gives body to the paint, but being transparent, detracts from its coloring or covering power. There is less white lead in the mixture to saponify with the linseed-oil to form a lead soap that is quickly washed away by storms. The oil also dries out and the white lead is rendered liable to chalk. Such coatings cost less than pure white lead, are more bulky, and by a little extra labor on the painter's part can be made to cover more surface than white lead and zine exide, and they save oil.

Zinc oxide 50 per cent, white lead 25 per cent, and Blanc Fixe 25 per cent, also stand southern and seacoast exposures better than white lead alone. When the above mixtures are used on wooden structures the barytes and silica act as fillers in the first cont, and the percentage of either substance can be greater than in the other coats.

Pure zinc oxide is a pure white pigment but little affected by sulphur fumes, and does not yellow the oil with which it is ground or mixed. It is of itself a good drier, and is used in the preparation of kettle-boiled oil. When it forms the principal pigment in a paint, other driers are detrimental, as the coating has a tendency to harden upon the surface only, and remain viscid below and peel readily. cause there is less lead in it and the atoms of each are better coated with the oil from the larger quantity of it necessary with the zine pigment. Mixed-zine oxide and white-lead coatings are, in general, more durable than either coating alone, provided both the pigments are pure. As zine oxide costs more than white lead, the adulteration of it is quite as general, and with the same substances. (See Chapter VI.)

It deteriorates by long keeping and loses much of its covering power, which can be restored by heating the pigment. When freshly made or heated and exposed to moist air, it changes, by absorbing earbonic acid, into the carbonate of zine (ZnCO₃). Painters say "it spoils," and grard against this change by closely covering or sealing it, or mixing it at once into a paste or paint, where the oil protects it, except those particles that lie upon the surface. The carbonate of zine so formed is crystalline, loses density, and hardens so that it cannot be pulverized or ground without extreme effort.

In this change S1 parts of zine oxide (ZnO), composed of zine 80.344 per cent, oxygen 19.656 per cent, specific gravity 5.42, corresponding volume 14.9, are changed to equal 125 parts of the carbonate of zine (ZnCO₃), composed of zine 52.153 per cent, oxygen 38.278 per cent, carbon 9.569 per cent, specific gravity 4.44, volume 28.1,

This chemical change, attended by so large an increase in volume (nearly double), if it takes place after the oxide has been spread as a paint, the whole coating will be loosened, and the loose particles will be carried away by storms; the action being similar to that which would occur should the sand mixed in the mortar of a plastered wall, when dried, change its volume to the same degree.

Zinc oxide is therefore not a permanent paint for open-air exposures, but for interior use is permanent; for though carbonic acid is present, the moisture is absent, both elements being essential for the change to a carbonate. Dry, gaseous carbonic acid does not affect dry zinc oxide. It is mixed with white lead for interior use to lessen the tendency of the white lead to darken by absorbing the sulphuretted hydrogen present in nearly all locations, a small

Zinc oxide is a hazardous pigment to use for external exposures when mixed with iron oxide, lead, or other color pigments. No process of mixing them can so associate the several pigments, even when ground in the oil, as to enable any one particle of either substance to thoroughly protect any particle of the others present, from the changes mentioned. Atmospheric moisture and goses will sooner or later reach the zinc oxide, whether in the first or other coats of the paint, and the inexorable laws of chemical change will govern the durability of the coating, the ultimate decomposition of which will

be determined simply by the amount of zine oxide present.

Zine sulphate (ZnSO₄) is a white pigment, and is often produced in the manufacture of zine oxide, the color of which is not affected by its presence, even if the sulphate is added to the oxide afterward. Zine sulphate has recently been brought forward as a desirable pigment for ferric as well as for wooden surfaces. In Germany and England it is used largely in all mixed paints, and has thus far proved to be very resistant to atmospheric influences in damp locations. Like zine oxide and the lead pigments, its cost is a great factor against its more extended use. The commercial zine-sulphate paints are adulterated with barytes, the natural composition of which is favor-

able for its admixture with the sulphate.

Zinc sulphide is a pigment introduced by Mr. J. B. Orr, of England. Its composition is, barium sulphate 70,50 per cent and zinc sulphide 29,50 per cent, the reactions being, BaS | ZnSO₄ | BaSO₄ | ZnS. The dense white precipitate formed is highly heated, then quenched in water, and finely ground and dried. Becton white, Olemn white, Orr's white, etc., are zinc-sulphide pigments. Great purity of the raw materials is required to produce a purely white product.

Zinc sulphide is largely used in the manufacture of enamel paints, linoleum, table oilcloths, etc. It does not continue to exidize after mixing with linseed-oil, as do lead exides, and can be considered as a saturated or non-siccative compound. It does not combine with resin, and therefore will not saponify. Exposure affects it by blackening the paint if a lead drier has been used with the linsest-oil, or a lead pigment associated with the sulphide pigment. In these cases a lead sulphide is formed which is dark-colored.

Yellow Seal. The Green Seal consists of one part zine sulphate and two parts of barytes. The Red Seal, of one part zine sulphate and three parts of barytes. The specific gravity of the Red Seal is 4.2. The Blue and Yellow Seals contain some zine oxide with the sulphate, and a greater percentage of barytes, and are consequently deficient in covering power. The large amount of oil taken up by the sulphate and zine oxide is counteracted by the smaller quantity taken up by the barytes.

Barytes costs about one-tenth as much as zine oxide or zine sulphate, and affords every requisite to grade up the weight of zine paints, even when a liberal amount of whiting is present, as is too frequently the case with most mixed pastes or color paints. (See Tests of Paints.)

Green-scal lithopone approaches closely the best brands of French zinc oxide, and does not require so large an amount of thinners as the American brands of zinc white, and it works easier. It is unaffected by sulphurous gases, and does not turn yellow when thinned. It will blacken if exposed to the sun before it is dry. Oils or driers containing lead or copper salts turn lithopone gray; neither can it be used with other colors having a lead or copper base.

Griffith's zine white, a chloride or sulphate of zine, is precipitated from a soluble sulphide or chloride of sodium, barium, or calcium. No iron should be present. The precipitate is dried, and then calcined at a low or cherry heat, with careful stirring; raked out of the furnace and quenched in vats of cold water, then levigated and ground. It is an oxysulphide of zine; some sulphate of magnesia accompanies the pigment.

A commercial zine white that is only a sulphate of zine is made by precipitating the pigment, by the addition of a dilute solution of sulphuric acid, to an acetic- or nitric-acid solution of litharge. Wash and dry the precipitate thoroughly. The clear liquor can be used repeatedly. All of the metals associated with the zine in the litharge are dissolved by the nitric or acetic acids and precipitated with them as sulphates, and are inclined to blacken on atmospheric exposure. The zine pigments formed by the precipitation processes are not as durable or reliable as those formed by the oxidation or sublimation The United States Bureau of Construction specifies one part of white lead to three parts of zinc oxide for the paint used on wooden structures on the seaconst, and has lately abandoned the use of zinc-oxide pigments on ferric structures wherever located. House-painters use from 20 to 50 per cent of zinc oxide when they mix their own colors. The great percentage of zinc oxide in the commercial mixed-white paints and colors has been referred to in Chapter VI.

Adulterations of Zinc Oxide, and Tests.

Patent zinc white is a sulphide of zinc mixed with baryta or strontia. Fulton's zinc white is the sulphide of zinc and barytes. Charlton's zinc white is the same.

In the adulterations of zine oxide with baryta, barytic white, permanent white, Blane-Fixe, constant white, etc., all of these substances are artificial sulphates of baryta, and are less crystalline than the natural sulphate, and cover better. Pure zine oxide dissolves entirely in dilute sulphuric acid, leaving no residue. If carbonate of lime is present, it effervesces with nurriatic acid, and the amount of this action in a measure indicates the amount of that adulterant present.

Zinc oxide lacks weight when compared with white lead for paint mixtures. Barytes in its powdered form supplies this deficiency, but has poor covering power; it spreads well and saves oil. The floated barytes—a finer grade floated from the pulverized natural mineral—has better covering power than the ordinary brands of this pigment, simply because it is finer. Artificial barytes or "Blanc Fixe" frequently contains pulverized silica or white glass sand. All of these substances are adulterants, and add nothing to the qualities of zine oxide except weight and a saving of oil that lessens the cost of the mixture to the manufacturer, but seldom to the consumer.

To test a mixed zine-oxide paste or paint for adulterations, repeated washings with benzine or other will remove the oil; then dry the residuum on blotting-paper. Dilute sulphuric acid completely dissolves zine oxide, leaving the adulteration or any other metallic base pigments unaffected.

^{* &}quot;United States Navy Yard Tosts of Marine Paints" Provident

has been ignited.

On a painted surface, slightly scratch the coating, and apply a drop of sodium sulphide of 100° Baumé. If lead pigments are present a discoloration will follow the application of the sodium.

CHAPTER VIII.

LAMPBLACK.

THE carbon group of pigments comprises lampblack, mineral or natural asphaltum, artificial asphalt, coal-tar, and graphite, either alone or as a component part of the paint.

Lampblack is the fine deposit or soot formed by the imperfect combustion of oil or fatty substances. Its composition varies greatly, depending upon the nature of the substance consumed in its formation, and the care exercised in the combustion process. Fatty oils and grease yield the best lampblack. Cond-tar yields a black of a brownish huc and is inclined to be oily. Resin furnishes a good black. If the combustion is forced it carries along some of the free resinflakes, and yields a yellowish resinous black of an inferior quality, not always free from grit and dirt.

Gas-black, the soot product from the combustion of hydrocarbon fuel or illuminating coal-gas, differs in molecular structure from the fatty-oil blacks. The gas-black particles appear to have a star form, and are not as suitable for mixing with white lead or zine white for tints as the fatty-oil blacks, though their color is densely opaque. The fatty-oil lampblack is filament-formed, and incorporates with the oil and oxide pigments better than the star or flake formed blacks. Gas-black is also made from natural gas burned under revolving cylinders; the deposited soot being removed by scraping. With proper care the lampblack so formed is nearly pure carbon. In a paint coating it has a tendency to become brittle, crack, and thake off after a short time. This possibly results from using too much drier or turpentine in the vehicle, as of itself it is a slow-drying pigment, and adds no drying qualities to the vehicle. It seldom appears in the market as gas-black.

Ground soot appears as a lampblack under various trade names. It contains ammonia, sulphuric and pyroligueous acids, rain-

on roofing material and ferric surfaces. With galvanized iron or sheet zine, the zine is reduced to either an oxide, sulphate, or carbonate at the expense of the zine covering, leaving the iron exposed to the action of the elements which produce corrosion, that is the more active because of the galvanic couple of the different metals. (See Calvanizing, Chapter XVII.)

Spanish black, or cork-black, is made from the combustion gases of burning cork. It is a good lampblack in color and texture if proper care be taken in the process; but charred cork and ashes are too often present in the product for its good.

Irory-black is made from chips of elephants' tusks and other hard bones free from fat. It should have no hister, as that indicates the presence of unconsumed fatty matter. Its use is almost exclusively for the preparation of the finest blacks for carriage, decorative, and artists' colors. Its high cost debuts it for use in ferric contings,

Horn-black, or animal black, is almost identical with bone-black, but is generally in a more finely divided state. Animal refuse, allumen, gelatine, horn and hoof shavings, etc., are subjected to a dry distillation in a still or retort; the black carbo-mecous mass left is washed with water and powdered in a mill. It requires about one and a quarter its own weight of oil for a paste. The great quantity of oil left in the black as it comes from the still is the reason for its slow drying. It is a cold, mild black, and when not well burned has a brown tint, dries badly, and is used for printers' ink, blacking, etc., also for the cheaper grade of black varnishes and paints.

Bone-black, made from a poorer quality of bones than ivory-black, is a warm, reddish-brown black.

Drop-black is an ivory- or bone-black blued with Prussian blue.

Charcoal-black is a finely powdered beechwood charcoal, made in Sweden, and generally marketed as Swedish black. It is a pure black in color, but has less covering power than the fatty-oil blacks.

Blue-black, made from vine-stems, is a better quality of charcoal-

this black come from the condensed gases as soot in the calcination of the above substances. Many other process blacks are sold under the above names.

Almond-black is made from fruit stones, nuts, etc. It is an intense black, and has the same qualities as Frankfort black.

German black is made from the combustion gases of any resinous matter, which escape into a long flue, at the end of which is a woodlen or other fibre hood, that collects the deposited soot.

English black is collected from the flues of bituminous coke-ovens.

Russian black is made from the soot of resinous dead fir- or pinewood. It is liable to spontaneous combustion if left for a short time
moistened with oil.

Prussian black, Berlin black, ochre-black, coffee-black, carth-black, lake-black, paper-black, and manganese-black are all inferior qualities of lampblack made by some one of the many processes, and from the many substances capable of slow and imperfect or smoldering combustion. Their color and qualities are quite as divergent as their names; all dry slowly with uncertain results in color and lustre.

Graphite-black, or ship's black, is an impure lampblack mixed with an inferior quality of flake-graphite, and can be known by its metallic lustre.

Coal- or shale-blacks are generally pulverized slaty bituminous coal.

The trade adulterants of lampblack consist not only of those substances that in the process of manufacture are imperfectly carbonized or vaporized, but nearly every other light substance that is black and can be ground to the required fineness. The coarse sout and scales deposited in the chimneys and flues from the combustion of fatty-wood and soft-coal fires, coal-gas, mineral oil, shale, and asphaltum, coal-tar, etc., in the several processes of distillation or burning for other products, all contain ashes; also acetic, pyroligneous, and sulphuric acids, ammonia, and tar to a notable extent, that condense in the carbon-atoms, and materially affect the color and quality of the lampblack. These are not always removed in the subsequent calcination that all lampblacks require to form the prime pure

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only on the surface, and remains viscid underneath, and is prone to peel.

Authracite and bituminous coals are ground and marketed as pure lampblack. They contain from 8 to 12 per cent of ash, also from ½ to 2 per cent of sulphur, and absolutely have not a single quality to recommend their use except their low price. From the large quantity of worthless lampblack selected for the finishing conting of most of the ironwork in the New York Rapid Transit Subway, it might well receive a special trade-name as the "Subway or Turnel black."

Carbon-black appears in the market as hydrocarbon-black, American gas-black, satin-black, gloss-black, jet-black, silicate of carbon, etc.

To make a pure lampblack requires not only a proper material, but as careful attention to the combustion of it and the subsequent processes for its preparation as the manufacture of any other pigment.

Pure hampblack made from a fatty oil is so finely subdivided naturally, that it requires no grinding. It is only ground in the vehicle to scenre a more thorough incorporation than is possible by stirring it in. It is of an oily feel and nature, and in combination with a good oil forms a more elastic and closer-clinging coating than any other pigment.

It is chemically and electrically passive, non-hygroscopic, non-corrosive, and less affected by heat, light, and peeling than any other pigment.

Its life in a paint coating is in a great measure exempt from all atmospheric influences that cause the decay of a paint. Its elastic nature reduces the frictional element due to the beating of storms, while the oxidation or decomposition of organic matter in the dust and from other sources is almost nil. It remains in place until removed by friction or the destruction of the vehicle, and can be painted over without the expensive torch-burning or scraping so necessary with other pigment coatings.

In some form or degree of purity lampblack enters into all of the black varnishes, enamels, and trade paints that have any marked of the coating.

Many instances are on record where a single coat of lampblack, like that used for the lettering and symbols on the old cross-road and tavern sign-boards, that have been exposed for a century or more, are still uninjured, while the surrounding colors and in many cases the wooden surface of the sign have been worn away, leaving the carbon lettering in full relief.

The iron-link-chain suspension bridge over the Merrimae River at Newburyport, Mass., was made of Norway cold-blast iron, and creeted in 1810. It was painted with two coats of pure lampblack and raw linseed-oil over sixty years ago, and is still (1903) practically free from corrosion, though in an exposed position, subject to sea air and fog influences for days in succession.

The use of lampblack to delay the "setting" of red lead is fully described in the article on red lead. It does not, however, prevent the failure of red-lead coatings when they are exposed to the action of hydric-sulphide fumes, but is not itself affected by them.

A good test of the quality of a lampblack is to place the sample on a piece of blotting-paper and pour a little ether on it until the paper is soaked with the other, percolating through the black. If on the evaporation of the volatile and removal of the powder the under side of the paper appears fatty, the lampblack is of poor quality.

Animal charcoal and bone-black or ivory-black are strong bleaching agents, and it is possible for them to uncolor overlying contings. The oil protects them somewhat from this bleaching influence, but where long stability of color or lustre is required, it is better to use blacks not of an animal nature.

CHAPTER IX.

MINERAL OR NATURAL ASPHALTUM, ARTIFICIAL ASPHALT (WHICH INCLUDES COAL-TAR AND ITS PRODUCTS, PITCH, MINERAL WAXES, ETC.).

Mineral or natural asphaltum. There are a large number of these. known as Egyptian, Bermudez, Trinidad, Mexican, Cuban, Californian, Chinese, etc. They all vary greatly in character and purity, and are the residual products of petroleum when the light hydrocarbon elements have been evaporated by natural causes. They contain vegetable and mineral matter, subpluric and other acids that must be removed by boiling or distillation to render them suitable for enamels, varnishes, or paints. Asphaltum is not to be confounded with the product of coal-tar distillation, called "asphalt," which, having a certain resemblance to the natural asphaltum in some of its physical qualities, is chemically very unlike it. The name asphalt being enrelessly applied to both the natural and artificial or coul-tar product, naturally leads to some confusion on the subject, They are, in fact, so widely apart in all their essential qualities that they cannot be appropriately coupled together as relating to the same substance.

The characteristics of asphaltum used for ferric coatings are briefly given: Asphalt, bitumen, or mineral pitch, specific gravity 1 to 1.68, softens at 170° F, and melts at 212° F, (coal-tar asphalt softens at 115° F,). According to Boussingault (Am. Ch. Phys. [2], X1V, 141) it is a mixture of two definite substances, viz.: asphaltene, which is fixed and soluble in alcohol; and petrolene, which is oily and volatile. The greater part of the latter may be volatilized by distilling the asphalt with water. The chemical composition of bitumen is:

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per cent purer than Trinidad. Samples of Bermudez analyze 97.22 per cent of materials soluble in bisulphide of carbon. A large amount of these materials is also soluble in ether, showing that the bitumen contains large amounts of petrolene.

Petrolene in Bernudez 81.63. "Trinidad 80.01.

Egyptian asphalt is the purest of all the varieties of asphalt, but is not procurable at present in commercial quantities required for pavements or paints, but is used in the finer qualities of japanned or enamelled wares, baked coatings, varnishes, etc. Samples of it frequently analyze 99.5 per cent of soluble matter.

Asphaltum yields by dry distillation a yellow oil, consisting of hydrocarbons mixed with a small quantity of oxidized matter. It begins to boil at 90° C., but gradually rises to 250° C., giving oils of specific gravity during the boiling, viz. from 90° C. to 200° C., specific gravity=0.817 (at 15° C.); that which boils between 200° C. and 250° C., specific gravity=0.868 (at 15° C.); both portions giving by analysis 87.5 carbon, 11.6 hydrogen, and 0.9 oxygen, which is nearly the composition of the oil of amber.

These asphaltum oils, treated with sulphuric acid and then washed with potash and subjected to dry distillation, yield a number of oils which are *insoluble* in water, or strong nitric acid, and are but little affected by strong sulphuric acid, but are very soluble in alcohol or ether.

Asphaltum has no metallic base, and can be classed as a gum or resin, hence but a small amount of it can be incorporated into an oil vehicle for use as a paint. Bisulphide of carbon and benzine usually form a large percentage of all vehicles in asphaltum paints.

The principal merit of some of these paints consists more in the name than the quality. If it is once considered that only about 10 per cent of asphaltum enters into the composition of the well-known street pavements, and that so little quantity as this amount, however it may govern the other constituents of the paving compound, has to be put in place or applied hot, and cannot be used or compounded in any other manner, it may be apparent that, notwithstanding the

market. Even with this small amount, and with the best of boiled or raw linseed-oil as the vehicle, the paint is difficult to dry without the use of strong metallic salts mixed with the oil to aid its oxidizing or drying quality; and if a quick-drying paint is wanted, these oxidizing materials are added in such amounts as to materially affect the life of the paint.

When the color of the paint is other than black or steely gray, it may be doubted if any asphalt will be found present under the closest analysis; and the red and brown colored samples will be found to rely almost whelly upon exide of iron as the base of the pigment, under whatever mane it may be masked.

Gilsonite, a mineral resin associated with natural asphaltum, is used largely as the principal pigment in these paints. Gilsonite, asphaltum, petraleum, cannel and bituminous coal and shale, all shade off into each other so gradually, and form so numerous a class of bituminous mineral substances, that it is difficult to determine their exact relations. The fluid elements of the hydrocarbons evaporate, and as the heavier portions solidify, they oxidize with a loss of hydrogen, and change until over a hundred different bituminous mineral substances can be determined from the hydrocarbon group.

The general composition of the numerous class of petroleums, after the evaporation of the lightest hydrocarbons by nature in the form of natural gas, is, viz.:

Cride Oil 26° Banné - Djotillates. Commercial Names.	Approx mone Degree Haitme	Specific Gravity.	Weight of 1 1' S Gall in Pounds,	Percent- age Oh- tuned. Approx- mate.
Cossoline. Renzine. Kerosene (filuminating-oil). Heavy kerosene (mineral sperm). Clas distillate. Light lobricating (spindle-oil). Neutral oil. Heavy lubricating-oil. Valve lubricating-oil. Asphalt Cende), containing 4 to 7 per) cent of sulphur.	75 76 63 45 38 40 28 26 23 21 14 15	0 6820 0 7253 0 80 0 8333 0 8866 0 8074 0 915 0 9271 0 9655 1 to 1 60	5 69 6 04 6 06 6 94 7 38 7 48 7 62 7 72 8 04 8 344 to 13 350	3 5 4 6 13 15 8 56 10 18 8 10 10 12 5 6 4 5 11-12 5 to 13

Coal-tar is a generic term applied to those bitumens which are extracted during the destructive distillation of bitundinous coal for gas or coke. Commercially, the name is also applied to water gas tar. The nature of the tar varies with the nature of the coal, and with the processes employed in its production as a waste product in the manufacture of gas or hard coke.

There is no known method of describing accurately the true composition of the coal-tars. No two are identical in every respect, although many are identical in every essential respect. Variations also occur from the admixture with the coal in process of distillation, of greater or less quantities of oils of various kinds, used for the purpose of enriching the gas. The tars vary in the amount of bitumen they contain within the limits of 60 to 92 per cent; also vary largely in the percentages of oil which they contain, and in the quality of the The non-bituminous matter in the tar is generally carbon. which is synonymous with lampblack, and was, of course, a hydrocarbon before the hydrogen was eliminated by combistion. Conf. tar cement, or asphalt, is a residue from the distillation of coal tar-Its hardness or flexibility is due to the percentage of the oil left in it. and may vary from 16 per cent in one quality of coal tar to 52 per cent in another. One per cent of oil taken from one coul far will produce a greater hardening effect than 14 per cent taken from another tar, and the degree of heat necessary for distilling off the oil may vary from 200° to 600° F., even when supplemented by mechanical agitation, or by blowing superheated steam or air into the still during the distillation process,

The average analyses of a large number of samples of coal tar from coal-gas retorts gave for a 40-gallon barrel, specific gravity 1418-1.10:

 $1\frac{1}{2}$ gals., or 3.75%, of light oils, consisting of heuzole, implithat and carbolic acide $9\frac{1}{2}$ gals., or 23.75%, of heavy oil, consisting of crossote-oil and anthractic, etc. 29 gals., or 72.5%, pitch.

Boiled in open kettles, this tar should be reduced from 15 per cent to 25 per cent, according to the duty required of it. The tar resulting from the distillation of petroleum oils for water-gas is of a decidedly inferior quality to that obtained from gas-coals, and is

better adapted for coating the cruder forms of wood constructions, piles, dock-timbers, fence-posts in the ground part, than metal-work. But this same oil-tar, if distilled at heats from 600° to 800° F., forms a pitch of almost adminantine hardness when cold, and resists almost all corrosive agents and solvents except those of the hydrocarbon class.

Analysis of a by-product coke-oven tar:

Naphthalene	12,00	per cent	
Anthracine	0.30		
Tar acids	7.00	16 66	
Tar bases	1.60	11 11	
Water	2.00	11 11	
Pitch	77.10	11 (1	
	100 00	() ()	

100 00 " "

When the concentration of the gas coal-tar is carried to the 25 per cent or 30 per cent stage, the product is comparatively odorless, or at least is not any more objectionable than that from oil paint, the pangency due to the light oils and carbolic acid being dissipated. In the distillation of coal-tar, until the final residuum of coke is reached in the still, there are no constituent oils derived from the process that do not gradually volatilize by the heat of the sun or approximating temperatures; and all coal-tar or hydrocarbon products suitable for use in, or as paints, also become fluid when exposed to heat; in fact, but few of them are applied in any other condition than while hot. They are all liable to run on vertical or slightly inclined surfaces, until by evaporation they are so advanced on the road to brittleness that they solidify, and by a little further progress in the same direction they become brittle and scale off on the least mechanical disturbance.

In the production of an ordinary standard roofing-pitch from a coke-oven tar, the distillation ran, viz.:

150° to 170° F.					
Analysis of a standard coal-gas tar (specif	ie gr	n ít	<u>y</u> 1.	21).	
Carbon	. 89	21	Int	cent	
Hydrogen	. 1	95	**	••	
Nitrogeu.	1	05	• •	**	
()xygen		13:1	**	**	
Ash	. (rnee			
Volatile sulphur	. 1	56	**	**	
	1 ()() (H:		*1	
In combustion it gave British thermal un power from and at 212° F.—16.4 pounds of v Analysis of water-gas tar from gas-oil (sp	vater				
Carbon					
Hydrogen		1 1:1			•
Nitrogen) 11		11	
Oxygen		3 65		**	
Ash		raci			
Volatile sulphur		11 :17		**	
	10	1) (8	,	**	
In combustion it gave British thermal m	iis 1	7.19	13.	Exa	morativ
power from and at 212° F. 17.8 pounds of	wate	41.			,
In the distillation of an oil water-gas to	ir by	111	Jo	lin I	· Wan
the products obtained at the several stage					
follows (specific gravity of the crude tur 13.5)					
Distillation heat, F.					destillate
240° F. water				gran Jark	
			25	1	•••
9409 10 hold off			200 1 4		
240° F. light oil		- (1	.41	**	
240° to 336°			Dč. 1	••	**
240° to 336°			()()		**
240° to 336°		. 3 :20	(H)	••	
240° to 336°	• • • • • • • • • • • • • • • • • • • •		(H)	••	**

At 400° F, the distillate became heavier than water. The residues obtained at temperatures of 550° to 617° F, were soft pitch, but would not flow. There were from 12 to 15 per cent of free carbon in the oilgas tar, while 5 to 8 per cent are the usual amounts found in an Otto-Hoffman coke-oven coal-tar.

The acids and ammonia salts in crude coal-tar must be climinated by boiling or distillation when used for coating ferric bodies. If they are not removed, the tar, either hot or cold, is one of the most unreliable and unmanageable of coatings. (See Dr. Angus Smith's experience, Chapter X11.)

The mineral waxes derived from coal-tar are the most reliable of all the coal-tar paint products. They are especially not affected by "sweating." They are an intermediary substance between the fluid and volatile elements and the heavy ones; and retain some of the volatile element that, as it slowly evaporates, causes the paraffin to crack badly and change its volume. The spaces between the tension-chord and other eye-bars in modern bridge constructions, lying so closely together as to be incapable of inspection or repainting to protect them from corrosion, are often filled in with melted paraffin as a protection from rust. It requires but a short period for the wax to harden, shrink, and crack, and expose the ferric bars. As well expect a cracked varnish coating to protect the surface it covers, as one of cracked paraffin.

CHAPTER X.

ASPHALTUM PAINTS AND CARBON VARNISHES.

Asphaltum paints are proprietary products, and vary in composition and quality quite as much as does the substance from which they derive their name. There is no standard of excellence in asphaltum paints.

A small amount of some quality of mineral asphaltum or gilsonite, mixed with varying amounts and qualities of the trade lampblacks, constitutes the pigment for the numerous brands of quick-drying paints used to blacken a large class of ferric bodies that need a coating for appearance rather than protection from corresion. The catchy name often secures their use on more important structures, where the price at which they are offered should promptly condemn them before trial.

A supposed better class of asphaltum paints or so called varnishes, similar to the "Maltha," "P. & B.," and other trade-mark designations, are freely marketed as superior paint products. They are in no sense varnishes, but simply the above mentioned class of pigment substances, mixed with bisulphide of earlier, bearing, and other uncertain hydrocarbon liquids and oils, the latter often containing more resin-oil than linseed-oil. They are not compounded by heat, as all true varnishes are. They have had an extended trial for over fifty years on important ferric structures, - naval, hydraulic, and other work, only to fail after a brief exposure. Wherever placed in competition with other earbon or metallic-base contings they are invariably found low in the column of merit. As a rule they operad easily and show well at first, but when the volatiles have evaporated, especially if they have been subjected to a moderate heat test 140° to 180° F., they become brittle, turn brown, crumble, and are easily removed. The application of these paints, containing bisulphide of carbon, is attended with extreme danger from fire, even on external exposures. The vapor of bisulphide is very explosive at low temperatures, also disastrously injurious to the painters or others breath-



Fig. 20. Animi Fossil Resin

An account of its application to water-mains, where it resulted in the insanity and death of a number of the painters and workmen engaged in painting and laying the pipes; also in the utter failure of the coating to protect the same pipes from corrosion, is given in "Transactions American Society Mechanical Engineers," Vol. XVI, 1895, Paper 637. Also in Engineering News, Feb. 7, 1895, and April 4, 1895.

A further demonstration of the inferiority of these asphaltum paints in competition with other oil paints and black varnishes is given in a series of tests made by Mr. Max Toltz, C.E. The Report was read before the Society of Civil Engineers, St. Paul, Minn., and reported in the Journal of the Association of Engineering Societies, 1897. It was also briefly referred to in "Transactions American Society Mechanical Engineers," May, 1901. (See also Bisulphide of Carbon, Chapter XX.)

Asphaltum varnishes or carbon paints in which the vehicle is practically a linseed-oil varnish, compounded by heat, and of the same nature as a baked-japan vehicle in which the carbon-blacks and other pigments are ground, are very reliable for protective coatings. They seldom fail under the severest tests of marine or other corrosive exposures.

Fossil Resins.

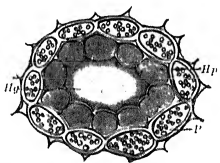


Fig. 10. Section through a resin passage of Abies excelse (fir and sprace-trees). The cavity Hq. as well as the thin-walled cells Hp, are filled with semi-fluid resin. The thick-walled cells P contain starch.

Fossil gums or resins, under the general name of Copals, are those

manufacture, many of them possessing pectant quantities. The first are the fossil resins found in the beds of rivers or in the earth where they have lain for centuries. The hardness of these fossil gums appears to depend upon their age and the pressure that they have undergone while buried. Amber is the hardest and most valuable of all resins. Only the refuse of black amber is used for varnish. Amber varnish merely means amber-colored varnish. There is no amber in the commercial brands.

Copal is the next in hardness; it comes from Zanzibar, and is known in the English trade as "Animi," from the insects embedded in it. Being very difficult to dissolve, it is distilled until it loses from 20 to 25 per cent, when it can be dissolved in boiling oil. There are three varieties of it, and many grades.

"Animi" is now the technical name for the South-American copal, and comes from Brazil.

Sierra Leone copal has nothing to do with Sierra Leone except for its name. It comes from the river-beds in the interior of Africa. It is the only African resin that will dissolve in cold alcohol. Its color is not as good as the Zanzibar or best Kauri, but it is harder than the Kauri. It is mixed with the Zanzibar for hardness, itself giving toughness to other fossil resins.

Other African copals are the Pebble or Pebble-stone—which is the hardest—Acora, Loango, Gaboon, Congo, Benguela, and three sorts of Angola.

Manila is of two kinds,—a hard and a soft; neither are fossil gums. They come from the Philippine and other islands, Borneo, Singapore, etc. This gum can be used as it comes from the living tree like the crude resin from the American long-leaf pine.

Danmar is a recent resin from trees not extinct, and contains the most water. When it forms the principal resin in a varnish or varnish paint, it appears to be always drying, hence the danger to any other coating spread over it. It is the resin used with enamel paints to give the high gloss characteristic of these coatings.

Sandarach is a resin yielded by the barberry-trees of Northern Africa. It is used to a considerable extent as the basis of spirit varnishes.

Kauri or Courie from Now Zooland to the salesting to it

when it has lain in the earth for centuries it becomes hard and valuable. It is very indifferent to the action of sulphur gases, and is more colorless than the other fossil resins. It is easily dissolved, and melts more readily than mastic, but less so than the common resins. It is allied in composition to Dammar resin, and is from two to nine times cheaper than the other fossil resins whose prices range in the order of commercial quantities as follows:
$\begin{array}{cccccccccccccccccccccccccccccccccccc$

Fourth, Zanzibar, best	•
The general composition of all fossil or other resins is Their specific gravities at 60° F, are:	(' ₂₀ [] ₄₀ () ₂
Yellow-leaf pine-resin (dark colophony) 1.100 " " " (whitish opaque) 1.047 to 1 " " " " (yellow transparent). 1.084 " 1 Shellae, D. C. (dark colored) 1.123	1.083
" B. (blenched)), 905 L, 063
" Very old	. 054 . 067 . 062
Benzoin, Sium	. 145 . 165
Tolu, old and brittle	.236 .231 .074 .038 .056

Shellac is an animal resin produced from the banyan- or fig-tree

Kauri (Australian).....

1 115 " 1,050

1.082 " 1.018

lacca, exudes a sap that the insect transforms by digestion into a resinous excretion (lac), with which she enerusts her eggs and herself. The insect is indigenous to the forests of India. The exudation of the sap from the lac-tree is somewhat similar to that produced by an insect or parasitic fungus on a species of oak (the gall-oak) that produces the "gall-nut" used by dyers and in pharmacy. The lac, when sent to market, often contains the eggs of the insect, and is called "seed-lac."

The lac secretion is dissolved from the twigs and branches of the tree in hot water, the solution is then evaporated on hot revolving cylinders, or in shallow paus, then scraped off in the form of thin sheets, broken up, and forms the commercial shellar, graded generally as D. C. (dark colored), L. C. (light colored), B. (bleached), etc.

The coarse qualities of the melted hie, when dropped into rounded pieces 1 to 1½ inches in diameter, are called "button-lne," and when in larger pieces are known as sheet-lne or "piece-lne."

The best quality is kusum-lac, from the kusum tree (Schleicheratrijnga), which lasts about ten years after being stung. The twigs from this tree are of a light-golden color and furnish the orange shellae; coming principally from Siam. The second quality is furnished by the dhak or palas, from the Butea frondosa. The third quality is the pipal, from the Fiscus religiosa. All of the lac-trees except the kusum live only from two to three years after being stung by the insect. Commercial shellaes are extremely variable in quality and price. The best grade of fine Orange D. C. lac brings £10 12s, per ewt. in London; "Native Orange," £8 9s, per ewt.; "Garnet," £7 8s.; "Native leaf" and "Button," £3 8s, to £3 6s.

The composition of shellac is given by Mr. Halstend* as

	Stick-lac.		Bood lac.				Shellar.			
Resin. Coloring-matter Wax. Gluten. Extraneous and nitrog-	68.00 per 10.00 " 6.00 " 5.50 "	cent, u u	88. 2 4 2.6	50) 50)	44 44 144 1444	rent.	$\frac{0}{4}$	181 261 (R) (R)	* * *	cent.
enous substances	6.50 "	"	1.0		11	**	2	80	**	ti
Loss	4.00 **	The same	1.1			ee Talkanaan	1.	80	**	**

A more complete analysis by Dr. Johns* shows that 120 parts of stick-lac consist of

### Property of the Control of the C	
An odorless common resin	80,00 parts
A resin insoluble in ether	20.00 "
Coloring-untter analogous to coclineal	4.50 "
Bitter bulsamic matter	3.00 "
Dim-colored extract	0.50 "
Acia (laccie acid)	0.75 "
Katty matter like wax	3 00 0
Skins of the insect and coloring-matter (the latter furnishing	
food for the grub when intched)	2.50 "
Solts	1 25 "
Earths	
Loss	
4407011 1 1 1 1 1 1 1 1 1	0.70
	190 00 46
	120,00

Shellac dissolves readily in alcohol, benzine, muriatic and acetic acids, but not in concentrated sulphuric acid. It dries solely by the evaporation of the solvent, leaving the thin film unchanged, the only use of the solvent being to spread the varnish. When alcohol is used as the solvent, the varnish can be spread over damp surfaces, as the alcohol will take up the moisture without much apparent injury to the coating, though this will be longer in drying, as the water must be evaporated with the alcohol.

Shelhe can be applied to ferric surfaces, and in under-water (fresh) exposures it generally will remain about two years without any great deterioration. In salt water, however, it will not stand a week, and when exposed to the sun and air, will be destroyed in about a month.

Each of the fossil resins represents a class that have many varieties, but none of them are coniferous. The latter class are those that furnish the turpentine and common resins of the present day, which are of the least value of any of the resins for a straight varnish or a pigment varnish. Their use in a varnish is principally on account of their cheapness and the slightly improved brightness they confer.

Records of the protective nature of some of these varnish paints show that a suitable combination of linseed-oil and a resin is a better protective vehicle than oil alone, yet the *smaller* the proportion of the company class of resins, the more durable was the conting.

compositions and quality of the varnish required to meet the conditions to which it is to be subjected. Heats approximating the charring-point of the oil, 450°-500° F., are necessary for a thorough blending.

Varnishes and varnish paints dry better if moderately warm when applied, or if applied to a warm surface. Manufacturers of pianos and other highly finished surfaces on wood subject their work to 200° to 250° F, to aid the drying and to harden the coating.

But cheaper materials and processes than the above are employed to produce coatings to compete with the basic metal pigments for use on ferric bodies. This careless compounding has resulted in lowering both the price and quality of varnish paints, until many of the commercial varnishes fall below the average of the better class of straight pigment oil paints for protective coatings on ferric structures.

For trade convenience, 100 pounds of resin are taken for the unit of composition, and with this unit, 8, 10, 20, or any number of gallons of oil rated at 7.8 to 8 pounds per gallon, are compounded for the different grades of varnish, known as 8, 10, 12, etc., gallon varnishes. To designate the kind of resin used, the initial letter of the kind of resin that is employed is taken, viz.: An 8Z varnish means an 8 gallon Zanzibar; an 8M, an 8-gallon Manila, and so on, both for the single letters or with a combination of the letters.

In the color varnishes or so-called enamel or paint varnishes, where the pigments are ground in the selected brand of varnish employed for the vehicle, the designated letter of the resin in it is generally lost or withheld, except as specially furnished by the manufacturer.

All of these varnishes or paints are best thinned with turpentine to the proper consistency required for the brush. It is better for this purpose than oil. The heating of the oil and resin together for the varnish has so thoroughly incorporated them, that no free oil is present to exert any change or action in the drying process, separate from that present in the coating as a whole, and which the addition of free oil as a thinner would disturb.

Benzine, or other distilled hydrocarbon liquid, should never be used in the composition of varnish or varnish paint. Their quick evaporation results in making the coating porous, and lighter to

An essential point in either a straight or a pigment varnish is that the linseed-oil should be made from ripe seeds, cold pressed, and be well aged, and its "Mucossities" or non-drying elements (nearly 6 per cent of it) should be removed, in part at least, or so changed in character as not to be readily decomposed in the natural oxidation of the vehicle in the process of drying.

Dingles's Journal reports the experiments of Dr. Sace (Nuremberg) to ascertain the nature of different resins, viz.: Amber, copal, common resin, dammar, elemni, caramba wax, mastic, shellac, and sandarach. All of them were reducible to a powder form. Amber, elemni, mastic, shellac, and sandarach became pasty before melting, the others became liquid at once. Amber and dammar did not dissolve in alcohol. Copal became pasty, elemni and zaramba wax dissolved with difficulty, while common resin, mastic, shellac, and sandarach dissolved easily. Caustic soda dissolved shellac readily common resin partially, but had no influence on the other resins.

Oil of turpentine dissolved neither number nor shellne; it swelled copul, dissolved caramba wax, common resin, dammar, elemni, and sandarach easily, and mustic very readily.

Boiling linseed-oil had no effect on amber, earnmba wax, copal, elemni, or shellar, while sandarach dissolved slowly; common resin, dammar, and mastic dissolved easily in it.

Petroleum ether had no effect on amber, copul, and shelhe, and was a poor solvent for caramba wax, common resin, elenmi, and sandarach, and was a very good solvent for dammar and mastic.

Benzol dissolved common resin, dammar, and mastic very easily, elemni and sandarach to a limited extent, caramba wax more readily than elemni, but had no effect upon amber, copal, and shellac.

Though gums and resins are generally spoken of us belonging to the same class, they are distinguished from each other by the solubility of the gums in water and the insolubility of the resins in the same liquid. The gums are insoluble in alcohol, while the resins are soluble in it. The so-called gum-resins are soluble in both water and alcohol.

The Trades Lournal Review (London) Dog 4-14 1901 n 15

intermediate product between the gums and resins, which invariably consists of twelve molecules, affiliating with "linoxin," the highest oxidation of linseed-oil. Dr. Kronstein produces an artificial resin identical with fossil amber, both in color and hardness; also has advanced his theory and process by producing the soft resins and balsams.

CHAPTER XI.

BAKED-JAPAN COATINGS.

For special locations and ferric constructions, viz.: riveted-steel water-pipe lines, anchor- and eye-bars, lattice-trasses, posts and beams, covering- or buckle-plates walled in or buried in musonry, and inaccessible for inspection, repairs, or repainting, a special coating called "baked japan" is being tested in a number of locations, the most prominent of which is a number of miles of steel water-pipe mains, 30 to 50 inches in diameter, riveted into a continuous length.

The process of manufacture and composition of the japan is similar to the black-varnish products, but a larger quantity of asphaltum, gilsonite, and other cheaper grades of gums and resins replaces the finer qualities of fossil and other resins. It is applied by immersing a hot pipe or other article in a hot bath of the compound, and upon removal from the bath and draining, baking it for a regulated period in an oven or muffle kept at an even temperature of 350° to 500° F., according to the size of the object to be coated, the composition of the japan, and the service required of the coating.

It fills all small interstices in the object, is clastic, will follow without strain all changes in temperature of the body coated, is perfectly impervious to atmospheric influences, running water, brine, acid, and alkaline and sulphur solutions, that affect the *ordinary* oil-paint coatings. Its cost per square yard of coated surface is naturally much greater than any brush coating, and will vary according to the conditions of its application. Its durability or life may be anywhere from ten to fifty times that of the ordinary oil-paint coating exposed to the same influences.

The composition of such baked coatings (and there are scores of them in practical use) appears to be of less importance than their proper proportion, and the care used in their combination, application, and final baking. It is reasonable, however, to expect that a ences tending to produce any other change than is subordinate to the original one that dried the conting.

Baked-japan contings, from the nature of their ingredients, are

electrically passive, except to currents of high potential; hence it remains for time to determine whether the stray electric currents, now a fruitful source of electrolysis in all ferric bodies that lie in the pathway of their return to their place of generation, will not find the rows of rivets that unite the several sections of the underground water-pipe lines coated with baked japan, the points to concentrate the electrolytic energy for a rapid corrosion of the pipe system at thousands of points in each mile, instead of a hundred or so, exposed in the usual spigot and bell method of joining the pipes. The electrolytic action at the rivets will be hastened by the difference in potential between the rivets and the pipe-metal both of which are of different potential from the japan coating. The brush paint coating applied to the rivet-heads will afford but little if any protection against corrosion or electrolytic action, as they will take place underscath the coating, and will require but a small development of either before they east off the paint and have an easier field for their progress.

Another source of corrosion which these joints can resist but a short time is the action of the acid elements present in all carths. In the case of these water-pipe lines exposed for miles to a great number of strong electric currents, the ordinary rate of corrosion from earth and water will be intensified, as in the water-tower stand pipe case eited in Chapter XXXIV of this volume.

While the baked-japan conting of itself leaves but little if any room for improvement in the conting of water mains, it will surely be a source of future regret that a better method of joining the short sections of pipe into a continuous line was not adopted than the riveted joints thus far used.

If the brush or modified japan conting applied to the pipe circular seams is adequate for their permanent protection from corresion, why incur the expense of a baked coating for the body part of the pipe? If it is not a permanent protection for them, then, as a chain is no stronger than its weakest link, there must be a great number of weak links in this method of constructing and protecting water-nine

ence between a good and a bad plan of joint construction to be a factor in determining which to use. That a number of American cities have this bad joint is evidenced from trade catalogues and other illustrations of this method of constructing large water-supply pipelines.

The question has been raised as to whether the baking of the coating effects any further chemical union between the oil and the other constituents of the dip, other than that developed in the process of manufacture? It is probable that it does, as the baking temperature is materially higher than that in the process of manufacture, The pipe-coating material before baking is readily soluble in turpentine, but after baking is not softened by prolonged digestion in hat turpentine, and but indifferently in hot naphtha. The preliminary heating of the pipe before immersion in the hot-pipe dip assures its adhesion and impermeability, as the air and moisture are practically excluded and the preliminary bond of the coating to the metal is perfect. The evaporation of the volatiles in the japan dip in the process of baking is so quickly effected in the earlier stage of baking, that the liquid or fused mass of the resins readily replaces them and fills the interstices caused by their evaporation, and ensures a smooth imbroken surface to the coating altogether different from that of a dried paint.

The so-called japanned or enamelled coatings used on sewingmachines and many domestic machines and utensils are generally of that composition that will give the best appearance. They are not proof against corrosion under many exposures that would be resisted by a good varnish coating or an earthenware salt glaze. As a rule they chip easily, and corrosion once established in these spots, spreads rapidly beneath the enamel and flakes it off.

A properly made enamel is essentially a glaze, similar in composition and properties to glass, and has all of the advantages and disadvantages of that substance. It is melted at a high heat, 1200° to 1400° F., and adheres to the surface of metals perfectly. Enamels generally resist the action of acid solvents, but are brittle and easily chipped off.

"The lest baked japans are intermediate between enamels and

In hardness, baked Japan is intermediate network variata and gass, or harder than gypsum and nearly as hard as marble." *

Baked black japans are made from linseed-oil and asphalt as a base, mixed with more or less copal resins, usually kauri, and are thinned with turpentine. Like varnishes, the more linseed-oil they contain and the less driers (oxides of lead and manganese) the more durable they are; but to get them to bake hard at a comparatively low heat, the proportion of oil is frequently decreased as much as possible and the amount of driers increased, forming an inferior, brittle coating easily injured by a slight blow or rough handling.

Modern baked-japan water-pipe coatings are very similar in character and in their application to Dr. Angus Smith's anti-corrosive water-pipe coating, that forms the subject of the following chapter.

^{*&}quot;Paints, Varnishes, and Enamela." A. H. Sabin, M.S., New York, 1896

CHAPTER XII.

DR. ANGUS SMITH'S ANTI-CORROSIVE WATER-PIPE AND OTHER COATINGS.

This compound was originally applied by Dr. Smith in 1840, and patented in England in 1850, and was first used in America in 1858 upon some pipes imported from Glasgow. Dr. Smith's original formula is not definitely known. Mr. James P. Kirkwood's Report on the Brooklyn Water Works, published in 1858, gave the following formula for it, and it was used to some extent upon the pipes for those works; evidently satisfactorily, for Mr. Peter Milne, engineer in charge of the extension of the works, reports: "That 36-inch pipemains laid for 35 years were found to be in perfect condition externally, and but few tubercules or other deposits were found on the inside of pipes." The pipes had been conted by heating them in an open furnace to about 500° F., and then immersing them in a bath formed from conl-tar, as follows: *

Coal-tar was distilled until the naphtha was removed and the material deodorized and of the consistency of melted wax or a thick molasses. This process also eliminated most of the tarry acids, and necessarily required considerable time and care to effect. Five to 6 per cent and in some cases 8 per cent of pure raw linsced-oil was then added and stirred in well. The bath was made deep enough to receive the pipes when placed in it vertically. The pipes remained in the bath until they had cooled down to the same temperature, about 300° F., or about 30 minutes for a 20-inch-diameter pipe. Careful attention was given to the length of time the pipes were to remain in the bath. A less time than 30 minutes for a 20-inch pipe gave an unsatisfactory result. For pipes from 4 to 12 inches in diameter, 15 to 20 minutes' immersion appeared to be sufficient to get a reliable coating.

When the coal-tar was distilled to the consistency of mineral

^{* &}quot;Report in relation to Proposals made by various parties to protect the

pitch or bitumen, or when common resin or Burgundy pitch was mixed with it and used as a bath, the pipe coatings became hard and brittle when cold, and the bath material would not answer, even where the quantity of linseed-oil used in it was increased to 15 ar more per cent.

The preliminary heating of the pipes to 500° F, before immersion in the bath, after a short experience, was found to be prejudicial, and was abandoned. The combustion gases of the heating-furnace that were deposited on the pipes appeared to affect the bonding of the coating to the pipe-metal, and the pipes when removed from the bath were not satisfactory, and new specifications for coating them were adopted.

These specifications required the same preparation of the conl-tar for the bath as given above, and for it to be kept at a temperature of 300° F, during the period of dipping. As the material was continually deteriorating during the dipping process, fresh material was to be added frequently, and at least 8 per cent of linseed-oil, as near as could be guessed at, kept in the bath, or added with the fresh pitch. The bath was required to be occasionally entirely emptied of its contents and to be refilled with new material. The old material after a few days' use was found to be hard and brittle like contains pitch.

Every pipe was immersed cold, but not frosty, and was to remain in the bath until it had attained the temperature of the bath, 300° F. This period was about 30 minutes for the 20-inch pipe, as in the previous specification. It required a brisk fire to be unintained under the bath to overcome the cooling action of the cold pipe when immersed.

The presence on the pipe of moulding-sand, dirt, noisture, frost, or oil and grease of any kind, was found to be detrimental to the application of the coating, and their removal was necessary before dipping.

The royalty paid Dr. Smith for the use of his formula, although no United States patent was in effect, was 37\(\frac{1}{2}\) cents per ton of pipe.

The price paid the English pipe-founders for coating the pipes ordered from them by the Brooklyn Water Works was \$1.25 per ton, for the years 1858 to 1860. American pipe-founders' and contractors' price for Dr. Smith's coating was about \$3.00 per ton as against a plain asphalt coating of \$1.83 to \$2.25 per ton.

perature of the preliminary heating of the pipe before immersion varied from 200° to 700° F., and the proportion of ingredients and their composition was equally startling, as were also the attending results.

Mr. Chas. Harmony, Chief Engineer of the Louisville, Ky., Water Works, who experimented for a number of years with Dr. Smith's formula as given by Mr. Kirkwood, reports: That "some of the pipes so conted, after an exposure of from six to eighteen years, were in as perfect condition as when first laid; but it was an exception, not a rule. In a majority of cases the coating on the inside of the pipe was all gone, and upon the outside surfaces it had apparently been of no importance in prolonging the life of the pipe. The difficulty experienced was, that in the heating of the bath to the temperature of 300° F., the coal-tar, resin, and pitch compounds became ummanageable by approximating the condition of boiling and volatilization, and going everywhere except in the place it was wanted. The coating was thick and apparently unbroken, but exceedingly brittle, and would crack and scale off in the ordinary process of handling."

The tension of coal-tar and pitch at a temperature of 300° F, is hardly less than that of water at the same heat, or equal to about 53 pounds' pressure. To maintain such a temperature in the bath in open atmospheric pressure is impractical, and the composition becomes unmanageable.

Other engineers report that the pipes after twenty years of exposure were found to be free from corrosion, but the coating had lost its bond to the pipe, and evidently remained in place because corrosion or other causes had not developed enough energy to cast it off against the pressure of the surrounding earth.

In these and similar instances of failure, the results appear to have been more markedly against pipes cast in greensand instead of a dry sand or loam-mould, evidently because the thick, vitreous, or partly fused greensand coating carried so much air in its rough, sandy surface into the bath, that it could not escape through the

the tough Bower-Barff skin, to which it is naturally so closely attached as to require pickling to remove. It is the subsequent baking that the pipe receives that renders this process a success. The composition of the bath can be varied greatly without much detriment to the protective nature of the conting, if the baking process follows the bath.

The generally unfavorable results attendant on the use of Dr. Smith's formula without the baking process, and the care and cost of it, determined the present practice of the pipe founders, which is to place the pipes for a short time in an oven heated to 250° to 300° F., then immerse them in the bath of hot coal-tar and pitch, and then cool them in the open air.

This coating is one of appearance more than of a protective or an enduring nature, and is only applicable to water-pipes, as in gas pipes, so treated the solvent action of the hydrocarbon vapor soon removes the coating, and the joints draw and leak worse than with the uncoated surfaces.

The carcless and indifferent boiling of coal-tar, to free it from its many acid and other impurities, makes it a variable and tunsatisfactory coating. Lime, gypsum, and other mineral substances mixed and boiled with the coal-tar to neutralize the ammonia, acids, sulphur, etc., only render the tar more unreliable and unmanageable. The carcless heating of the pipes and bath, also the length of time the pipes are left in the bath, and the subsequent treatment of the pipes when removed, are all factors in the indifferent results obtained.

Unless great care is exercised the small pipes will be overheated and unequally coated and brushed off, inside and outside. The larger pipes, requiring a longer time to heat, from the mass of metal they contain, will be underheated in the oven and cool down the bath to a lower degree than is requisite for a reliable conting. The subsequent brushing of the coating, both inside and outside, during the first period of cooling (a matter of from 30 minutes to 2 hours), promotes its reliability.

All coal-tars or their compounds of whatever nature used as a bath, or applied with a brush to any surface, hot or cold, are subject to the law of fractional distillation; that is, that such a mixture during

for a time, and so on.

The low boiling- or evaporating-point of the lighter elements of coal-tar or petroleum products makes them very uncertain in their composition, as changes of temperature in the bath from 220° to 350° F, are frequently noted without any change in the character of them that the eye can detect.

The character of the buth composition changes so continuously and rapidly during the dip that frequent additions of fresh stock must be made. These necessarily cool the hath, change its composition, and irregular coatings ensue to that extent that an entirely new bath is necessary.

The use of linseed-oil with coal-tur for pipe contings, as usually applied at the pipe-foundries, is of very uncertain value. It causes the dip compound to froth to nearly double its volume, and renders the coating lumpy in appearance and uncertain in its bond to the pipe-metal. It requires some effort by continual stirring to incorporate it with the coal-tar and pitch, and it is always liable to separate from them and float upon the surface, froth, soften the coating, and delay its drying.

Dr. Augus Smith evidently used a number of formulæ for pipe contings that contained linseed-oil as one of the ingredients. A long line of careful experiments with the best of coal-tar, pitch, and linseed-oil carefully heated and applied, gave almost uniformly good coatings. Using the commercial grades of these substances and having the ordinary day laborer to compound and apply them, the result was necessarily inferior, so much so as to cause the abandonment of linseed-oil in coal-tar pipe coatings by modern founders. If, however, the truth were acknowledged, the present coal-tar pipe coating would be found to be living on the well-carned and deserved reputation of Dr. Smith's compound.

Dead Oil in Pipe Coatings.

That part of coal-tar obtained in the fractional distillation of the tar between the temperatures of 410° to 750° F., and which contains creosote and anthracine oils (see Analysis, Chapter IX), is used to keep the pipe dip at a standard quality. It evaporates by One part of dead oil to about seven parts of coal-tar increases

the proportion of the beavy oils in the tar dip from about 25 to 35 per cent, and appears to make the coatings more uniform and of a better character than where fresh tar is used to reinforce the bath.

Thick the gives thicker and more uniform contings than thin the, and fresh the requires a hotter pipe to take bond than does old the.

Crude gas coal-tar boiled from five to six hours becomes a soft solid at atmospheric temperatures. During the boiling the temperature remains at about 220° F, for about an hour, then rises to about 290°, stays there for a time, and finally rises to about 350 F. All of the naphtha is removed and the tar is deodorized and reduced to the consistency of very thick molasses. If to sixteen parts of this tar 1 per cent of bailed linsced-oil be added, no frothing occurs even at 400° F. The mixture is thick and does not harden well on light iron pipes about 4-inch thick. On heavy iron pipes an inch or more thick, the conting hardens without difficulty; in some cases becomes too hard, is brittle, and flakes off readily by mechanical injury when handled. Dead oil added to thin the mixture causes no frothing. The experiments show that linseed-oil could be used with success and advantage with partially refined coal gas tar, and also indicates that its application requires more intelligent care than the methods employed with the usual crude tar conting.

Experiments with a refined far containing dead oil show that as high as 8 per cent of boiled linseed-oil resulted favorably in solutive and hardness of the coating. In other instances, where from 1 to 8 per cent of raw linseed-oil was used instead of boiled oil, frothing occurred and a poor coating resulted, evidently due to the presence and evaporation of the water in the raw oil. There is about 5 per cent of water naturally held in combination with the best quality of raw linseed-oil made from ripe flaxseed, and nearly 8 per cent in the oil made from unripe seed. With many brands of commercial linseed-oil, 10 per cent additional of water is frequently incorporated by stirring it in with a puddle or passing it through a mixing mill. All such oils are likely to be made up from fish, resin, mineral or vegetable and animal oils with no linseed-oil of any quality in them, and all the

to allow of its use to the great extent that pipe-founders require for their work. Heavy roofing pitch alone will run in moderately warm weather and becomes too soft and sticky for a pipe covering, unless hid immediately after conting. This is impracticable, and in cold weather it is too hard and brittle for transportation or handling.

Nine parts of heavy roofing pitch with one part of boiled linseedoil give a thick glossy conting less brittle than pitch alone.

Two parts of boiled linseed-oil with the nine parts of the pitch give a coating more elastic and tough.

Three parts of boiled linseed-oil with nine parts of pitch, the coating is more bulky and less smooth than with the others, while with larger proportions of the linseed-oil the coating partakes of the churacter of a slow-drying paint and requires baking, which gives it a superior quality.

Coal-gas tar belonging to the class of pyrogenic (fire-formed) compounds is unstable at ordinary temperatures, and is continuously decomposing by the evaporation of its immy hydrocarbon elements, intil nothing but the hard friable pitch is left, which contains nearly all of the sulphur element in the coal that forms the base of the tar product. Asphaltim, also a pyrogenic product, formed by the slow evaporation or distillation of petroleum, decomposes upon exposure by reason of the oxidation of the sulphur element in it, but is more durable than the coal-tar residuum or pitch (asphalt).

Asphaltum and linseed oil contings do not harden well, unless a hard grade of asphaltum is used.

Water-pipe Dips and Coatings,

There are many pipe dips upon the market, some covered by patents of doubtful validity, others secret or proprietary compounds of doubtful utility. Some of these compounds appear as pipe dips, also as brush paints applicable for ferric constructions other than pipes. (See Paint Tests, Chapter XXIX.)

The P, and B. Pipe Dip is a patent dip; the principal ingredients are probably an asplinit and candle-tar pitch. The latter is a pitch

qualities it may possess when on, are more than offset by the dangers connected with its application.

The P. and B. "Ruberine" consists of "ruberoid" dissolved in

naphtha. "Ruberoid" consists of California asphaltum or maltha and candle-tar pitch digested and vulcanized with sulphur. "Ruberine" dries rapidly, is hard to spread smoothly, but gives an elastic or rubbery coating. See tests of paints, New York Elevated Railway Viaduct, for an example of its qualities.

Mineral Rubber Dip (or Rubber Coating) is a secret composition

Mineral Rubber Dip (or Rubber Coating) is a secret composition whose appearance indicates that it is largely asphaltume. It is rather duller in appearance than the ordinary coal far or asphalt mixture. The dip requires a temperature of about 400° F, to apply, and then it is almost impossible to get a smooth or neat appearing surface. As yet its protecting qualities have not been determined.

"Bitumastic" Products comprise an enamed to be applied in a

molten state to the metal. Bitumastic cement is used hot for the preservation of ships' bilges and frames, instead of the usual by draulic cement coatings, and also for the protection of water pipes. Butumustic solution has bitumen for its base. It is a brilliant black paint applied the same as other paints, and is probably similar in character and composition to "Smith's Durable Conting." It has been used to a considerable extent on steel water pipes and for the limited period of test in that service is favorably spoken of - It dries in 24 hours, is said to be unaffected by acidulous, alkaline, or brine solutions. If applied to the clean dry surface of the metal, does not crack or peel when alternately wet or dry, or expected continually to running water in penstocks, water-wheels, etc. It is not affected by a moderate heat, nor by sulphur funces, and is furnished ready for use at \$1.75 per gallon. It is very volatile, and the packages must be well stirred while being used. Its covering power is about 400 square feet, and its weight about 9.5 pounds per gallon. "Crysolite" Enamel and Paint. "Crysolite" paint is made from

oil and a by-product, oven-coke. It weighs 9.5 pounds per gallon and spreads 500 square feet as furnished for a paint. When thuned with 12.5 per cent of oil, will cover 1000 square feet, and under general conditions in both hot and cold weather, dries completely in 30 hours.

ferric paints are). In the winter one-eighth of its volume of turpentine can be added to aid its spreading power, which can be made to cover from 800 to 1000 square feet. "Crysolite" paints cost about 75 cents per gallon mixed ready for summer use.

"Crysolite" coatings on annuouia tank-cars and reservoirs stand the action of ammonia liquors and gases better than most of the paints used for this purpose. "Crysolite" baked coatings under test resisted the action of carbomate of ammonia and ammonium chloride liquors for three months without injury. "Crysolite" under the influences of strong brine is more favorable than the commercial asphaltum or the ordinary coal-tar paints.

Hickenlooper's gas-pipe-dip compound, used by the Cincinnati

Hickenlooper's gas-pipe-dip compound, used by the Cincinnati Gas Light and Coke Company, the United Gas and Improvement Company, and several other gas companies, to coat their small service pipes, has a record of many years' exposure in the ground with few traces of corrosion. The failures thus far reported show that neither the process nor compound were at fault, but the lack of thoroughness and intelligence in its application. The pipes are first cleaned from rust and mill-scale and then immersed in the following dip and in the following manner. Twenty gallons of retort coal-gas tar are brought up to a boiling heat for a short time to evaporate as much of the water, acids, ammonia, etc., as possible, then 20 pounds of freshly slaked lime are sifted in from the top and well worked into the tar. Boil down to the consistency between a coal-tar and a pitch. When settled, add four pounds of tallow and one pound of powdered resin; stir until all are dissolved and thoroughly incorporated, then let the mass cool and settle; then ladle off into barrels. When ready for use, to each barrel of forty-five gallons of the above mixture add four pounds of crude india-rubber dissolved in turpentine to the consistency of thick cream. Heat the mixture to about 150° F, and immerse the pipe, previously heated to about the same temperature. After a few minutes' immersion the pipes are removed from the bath and laid upon skids to harden. The coating is somewhat softer than the usual pipe-founders' dip, and requires more time to harden, and continues hardening for a number of hours after cooling down to atmospheric conditions. The compound is especially

rough surface of the coating are the first to eatch the saline, sulphur, or other impurities in the water that form the basis for the development of the rust cones. The coating under these pits is the first to break down, being of inadequate thickness—probably only $z_{\rm tot}^{\rm tot}$ meh thick. The external surface of the pipe is as rough as the miside, and is not only exposed to the moisture to inaugurate corresion, but this moisture will contain all the acids in the scal or which the pipes are laid.

In all cases of the corrosion of water-pipes, it is the porosity of the coating that causes the formation of the tubercles and decay of the pipe. Nearly all of the dip contings, when tested by themselves or not in contact with ferric substances, were practically unarquired by acid solutions or running water.

In general, all pipe coatings, applied as they nearly always are in a careless, indifferent manner, will begin to show indications of tabereles in three years, and cases of tabereles in large pipes at the end of sixteen years have been noted, where the carrying capacity of the pipes had been reduced 20 per cent. Engineers noist carriestly take up this question of reduced enrying capacity of their waterpipes and decide whether it is not more economical to add from 5 to 10 per cent to the cost of the pipe in the form of better coating materials and better methods of their application than to submit to this decrease in flow, that always grows less with the age of the pipe, while the demand upon the service is always increasing.

Specifications for pipe contings appear to be of little use in producing a satisfactory coating, either in appearance or durability, as the directions they give are more often evaded than carried out by the foundry employés. After the pipes are conted and upon the drying skid, no ordinary inspection can determine the character of the coating other than its appearance to the eye or touch.

Testing pipe coatings is usually by the landmer to see whether the coating is so hard and brittle as to chip off in landling. The acid test determines the porosity of the coating by attacking the metal through the pures of the dip. A solution of one part muriatic acid and two parts water will affect both the coating and the covered metal mere at the end of sixty days than they would be affected by two ¹⁰ inch or more, the coating undermined and peeled off.
After all, in this age of specifications, inspections, scrimping, and adulterations, there is nothing equal to an honest and capable contractor, either for furnishing pipe, coating, inspecting, or laying it. Get such a one if possible and then watch him closely.

Generally, the time that the pipes are left in the hot bath does not exceed one minute, and is more often only one-half a minute. It is impossible to properly coat a pipe in one-half a minute, as the air carried into the thick turgid bath by the pipe will not escape in that time, and the top part of the inside of the pipe and the lower part of the outside of the pipe are uncertainly coated for this reason. The pipes are seldom turned over while in the bath, or outside while on the skids in the process of scraping and brushing off the surplus dip.

On pipes that are left in the bath for five minutes the coatings are markedly superior to those exposed for shorter periods. This is

on pipes that are left in the bath for five migates the contings are markedly superior to those exposed for shorter periods. This is the case whatever the nature of the conting, and is one reason why the Angus Smith and other older-day coatings gave such superior results to those coated by modern methods. They never had less than five minutes in the bath, and were often left for fifteen or even more in case of large pipes one inch or more in thickness. Modern pipe-foundry management allows no such exposures.

more in case of large pipes one inch or more in thickness. Modern pipe-foundry management allows no such exposures.

A coal-gas tar paint that has given very good results in the coating of gas-holder tanks and other situations where the metal is exposed to ammonia and sulphurous acids in solution and to alternate melting and drying under a great range of temperature, is made as follows: Coal-gas tar is well boiled to evaporate the water and light hydrocarbon elements and then 20 to 25 per cent of caustic quicklime is sifted and well stirred in to neutralize the acid elements in the tar. This is to be kept hot for a few hours and then an equal quantity of good Portland or hydraulic cement is sifted and stirred in thoroughly. The mixture is applied hot to the clean dry iron, and can be repeated soon as cool or dry if the exposure conditions are to be very severe. In the latter case, a little more cement should be added, so that the caustic lime and cement mixture will contain 50 per cent of each. The pigments thicken the coal-tar and prevent it from run-

ning under sun temperatures and give a bond to the brush coating of

The city of Perth, Scotland, where very pure water is obtained from the Tay, had their water-pipes coated with a solution of india-After 25 years of use every pipe under 5 inches in diameter rubber.

Derrick 1(oist Ambialt Ground 13 Steam

tank.

had been completely closed by corresion. many cases where the ordinary coal tar dip had been used on the water-pipes it scaled off in strips and was discharged at the house service-taps.

A pipe-dipping tank being required for some steel riveted pipes, 16 to 30 inches in diameter and 28 feet long, was exten porized from old material in the contractor's vard. and is shown by the following Fig. 21*

An old boiler-shell 3 feet or more in diameter and 26 feet long was fitted with a slightly dished wrought iron flange 2 feet or more in width all around, riveted to the top end of the shell. This served as a working platform, also to eateh and return the drip. The other end or lower one of the shell was riveted and caulked to a cast from plate head which carried on its inside face a concentric flange in the centre, to which was riveted steam tight a wrought iron pipe nearly as long as the, outer boiler shell. This inside pipe was closed stemm-tight by a conical head that also served as guide for the pipe when it entered the bath of pipe dip. The bottom flange was tapped for steam- and drain-pipe connections, which were fitted with the usual gates, worked from Fig. 21.—A pipe-dipping the surface of the ground, in or on which the tank was erected. The anumber space be-

tween the centre pipe and shell was filled with the coal far or other pipe dip to be applied to the pipe, which was kept hot by the steam

^{*} Engineering Record, Vol. XXXV, May 8, 1897, p. 489.

coated kept the bath mixture well stirred up and ensured a nearly uniform quality of its ingredients. It is obvious that this comparatively inexpensive device is adaptable for many occasions that would not warrant a more expensive plant.

A larger shell could be fitted with a number of the conicalheaded pipes with their separate steam- and drain-pipe connections and be available for dipping a number of pipes at the same time, and would certainly ensure a more reliable conting than where the pipes are immersed in a long horizontal tank.

APPROXIMATE RELATIVE COST OF VARIOUS PIPE-DIPS AND COATINGS.*

Couting.	Approximate Amount Ro quoted to Cont One 48- inch Pipe, 12 Ft Long,	Approximate Prices.	Cost of Material for One 48-inch Pipe, Ap- proximately 325 Sq. Ft, of Surface.
Crude tar. Pitch. Pitch and linserd-oil. P. & B. dip. Mineral dip. P. & B. universal paint. P. & B. rulerine. Tar varnish. Datch varnish. Saldn's baked japan.	14 gals. 14 " 14 "	\$3.00 per bld, (52 gals.) 5.00 " " " 75.00 per ton 75.00 " " 1.00 per gal, 1.00 " " 0.10 " " 1.75 " "	\$0.22 a 0.50 .70 .45 .75 1.50 b 1.50 b .15 b .40 b 2.60 c

- a. About 30 per cent of this was lost by evaporation,
- b. Estimated cost of this conting as applied with a brush. The wastage would be excessive as a dip, but the dip is the only practical way for its use on a large scale, hence the figures are not strictly correct.
- c. This conting requires a comparatively expensive plant and considerable skilled labor, which would largely increase the total cost.
- ** The Manufacture and Inspection of Cast iron Pipes, Thos. II, Wiggins, C.E., Boston. Civil Engineers' Association Journal, 1899.

CHAPTER NIII.

GRAPHITE AND GRAPHITE PAINTS.

Carbon assumes in nature three allotropic forms, viz.: Diamond, graphite, and amorphous carbon. Graphite itself assumes different forms, some of which are amorphous and others strictly crystalline in character.

If the three allotropic forms of earlien had each a characteristic name, no confusion would be liable to arise in speaking of them. We speak of the diamond and of graphite, and each is clearly defined.

In speaking of the third form we are limited to amorphous earbon. This form is found in certain stages which are not strictly amorphous or granular in character. Cake, for instance, is one form; the others are the unineral graphite-earbon or graphite, termed foliated (flake), amorphous (granular), etc. Graphite is found in many parts of the world and is of various degrees of purity, ranging from 60 to over 90 per cent of graphitic earbon in the foliated form and 20 to 60 per cent in the other forms.

The foliated is a designation for the thicker flakes in the Ceylon and like varieties, while flake is used to designate the thin flakes of the purest brands, similar to the Tieonderoga mine product.

The German (Bavarian), Siberian, Mexican, and some American varieties are amorphous and vary greatly in the amount of carbon in their composition, as will be seen from the following analyses:

The purest brands (Ticonderoga mine) have a specific gravity of 1.21 to 1.4. The amorphous varieties range from 1.80 to 2.25 to 2.79. When pure it is perfectly opaque, iron-black or steel gray in color, with a metallic lustre. Its hardness varies from 1 to 2, and it conducts electricity nearly as well as the metals.

Pure graphite or minerals high in graphite-carbon grind and feel greasy, and are repellent to moisture and oil. Flake-graphite above 80 per cent in purity, by long trituration with water, gat by reduced

against 1 to 2 for graphite when it contains 95 to 99 per cent of earbon. The Ceylon, Cumberland, Indian, and American flake varieties are the purest in earbon, and are used for pencils, crucibles, lubricants, stove-polish, foundry facings, etc., and to tone up the poorer varieties for many purposes.

Foliated graphites, though used for pigments, are not as satisfactory (for reasons given hereafter) as the amorphous variety, that, less rich in carbon, contains other mineral substances, non-corrosive, non-absorbent of atmospheric moisture and gases, either as individual substances or collectively as a natural mineral compound. That this feature may be duly considered when a graphite pigment is to be selected for a ferric structure the following analyses of amorphous graphite from a number of widely separated mines are given:

ANALYSES OF AMORPHOUS GRAPHITE

Siberian and German Muce.	U.S.A. Mines.
Por Cont.	Por Cont.
Graphitic carbon	28,39 to 33,48
Silien as SiO ₂	46.97 " 37.54
Iron soluble as $\text{Fe}_{\text{W}}O_{\text{H}}$	4,22 " 4.25
Alumina as Al ₂ O ₄	16.90 " 12.35
(Calcium as CaO	0.99 " 1.02
Carbon dioxide, water combined, so- dium compounds, iron pyrites, vols-	D #0 (/ T A)
tile matter and loss 4.09 " 8.22	2.58 " 1.36
Specific gravities, 2.25 to 2.79. Color, gray or d	rab. Hardness,

1.5 to 1.8. Fracture, granular.

Graphite from the Wisconsin mines analyzes, viz.:

Graphitic carbon	72.00	to	74.00	per	cent
Iron oxide	7.10	"	14,00	u	44
Silien	10.00	**	12,00	"	44
Alumina	8.00	ŧ	traces		
Water and undetermined	2,90	41	11		

The Mexican graphites are amorphous in character, are high in carbon, and have had but a limited use for pigments. When containing about 80 per cent of carbon they are better suited for lubricants

or vehicle, and are nearly as unoxidizable from moisture, atmospheric influences, combustion, and other gases as pure carbon. They are of an agreeable color and good covering power, and they work well in combination with other pigments; flow, bold, or carry well in the oil, and are as easily brushed out to cover as moreh surface as any good paint. They are not repellent to the oil, do not separate from it, nor set in the paint pot or barrel on long storage, either as a paste or paint. They are wholly self-supporting as pigments, contain no elemental substances that tend to reduce them to a lower plane by oxidation or slacking in the presence of moisture and gases. They require no body stuffing, either to bond them, or to keep them quiet, or from curdling or crawling during or after application, and they contain neither acids nor sulphur.

They are entirely different in character and composition from

the so-called silien graphites of commerce, many of which resemble carbonaceous sebists or inpure soapstone, or are compounded from flake-graphite and mineral substances of dissimilar character, such as barytes, silica, furnace slag, etc. These several substances, even if they are non-corrosive, or electrically or chemically passive of themselves or collectively, when assembled in a paint cannot be as reliable as are the same substances incorporated together by the processes of nature, each and every particle of which is of the same physical and chemical composition and equally affected by the vehicle, atmosphere, or other conditions that affect a paint.

They have not the merit of being synthetical compounds. No human care in the mechanical processes of grinding and mixing them, as a compound pigment or paint, can arrange them in sequence or in other than a haphazard manner.

Silica graphite paint is of a dark, lifeless brown; not objectionable on the enclosed ironwork of a building, but decidedly so for more prominent positions. Hence it is toned up by red lead or other twice pigments of agreeable color, but at an increased cost and a contributed element of danger in the disintegrating of the paint whenever hydric-sulphide fumes reach the red lead in the conting

Iron oxide is also used for toping effects, but the natural red hematite oxides are not strong enough in color to materially modify the dependence can be placed upon them to work together for a durable conting. An acid and an alkali will chemically form an innoxious whole, but this or similar action is dangerous in a drying paint and generally proves detrimental to the conting or covered surface.

High-carbon graphite is so easily adulterated with soapstone that if a pound of it be ground with three pounds of soapstone (specific gravity 2.7), neither the eye nor touch can detect the adulteration; only analysis will show it.

Graphite is one of the lightest pigments. Its specific gravity ranges from 1.21 to 1.4 to 2.38, while zinc oxide is 5.42, asphaltum 1.4 to 1.8, barytes 4.5 to 4.7, silica 1.9 to 2.8, gypsum 2.15 to 2.35, iron oxide 4.7 to 5.4, whiting 2.2 to 2.8, red lead 9.07, white lead 6.43.

The natural drying of a linseed or varnish coating is in the form

of a closely woven web of a fine fabric. This shows plainly on a freshly dried or drying surface, and explains the reason why two or more coats are necessary to give a smooth foundation for the last or polishing coat. Each subsequent coat fills the interstices of that underneath it, each coat repairing the other's deficiencies, as many folds of

if the brighter red copperus oxides are used. Their strong sulphur element sets into action an antagonism between every element in the coating and delays the drying of the paint, making necessary excessive amounts of strong driers to counteract even a small percentage of it.

Graphite paints are neted for being slow driers and require a liberal use of driers to get a firm coating. This is more apparent with fluke-graphite; its flocculent form and oily nature prevent the vehicle from bonding it. There is a movement in the paint during the whole process and period of drying that even the sharper and more angular form of the silica or barytes added cannot wholly overcome. Furthermore, these substances bring their own peculiarities into the coating and forcibly demonstrate the unreliable character of all compound paints. The greater the number of substances in a paint the less

a fine muslin will in the aggregate make an adequate covering from heat or light.

Now, it is the function of a pigment to fill these cellular formations in the drying vehicle, or rather, while being applied with a brush, for the atoms of the pigment, mechanically arranged in brushing out

pigment and the vehicle.

If the pigment atom be splintered like a sliver of glass, or of only length and breadth like a flake, then the natural cellular formations in the drying vehicle cannot be realized. Such shaped pigments are arranged with the sharpest angles and edges upright to the drying surface, and are not well covered in or embedded in the oil, hence dry with a rough surface that will hold moisture and dust and quickly decompose and disintegrate them from their bed, when more moisture, cinders, and dust take their place and the cycle of action is repeated.

The rough character of all paint coatings containing siliea, barytes, furnace slag, etc., is distinctly apparent to the touch. A round numble does not bed itself in a cement as well as the cubical block from which it is made, neither does a beach-worn sand or a quick-sand atom, with the best of cement, make a good mortar for the same reason. The splinters, flakes, and round atoms are more easily removed from their beds than a square atom.

Both the amorphous and flake graphite pigments (not associated with foreign substances as adulterants) being electro-negative, are less affected by eatalytic or electrolytic action caused by the juxtaposition of electro-positive substances in the coating or surface covered, or by hydro-sulphide gases, than any class of pigments, lampblack alone excepted. This is a valuable feature in any paint, whether applied to iron or wooden bodies, and in the future will insure a more extended use of graphite paints instead of the iron oxides, and compounded or patent paints, to the curcless use of which most of the corrosion in progress upon important ferric structures is directly traceable.

As a general rule graphite uninerals that contain about 40 per cent of graphitic carbon have proven to be better for pigments than those richer in carbon, for the reasons given before, the principal one being that they are less repellent to the oil and bond better to it, and do not appear to be affected by combustion gases.

Amorphous graphite coating applied to boiler tubes exposed to internal firing and the action of hot water under pressure of eighty or more pounds per square inch for two years was uninjured and fresh as when it was first applied.

were replaced, each alternate tube was coated with Lake Superior graphite paint, the others being autocated. At the end of over two years a number of the tubes were removed and their condition ascertained. The unpainted ones were again covered with the hard, flinty scale, that required the use of a holler-scraper to remove. The painted tubes were covered with a light floculent coating of the scale, that could be brushed off with the fingers, showing the bright, clean

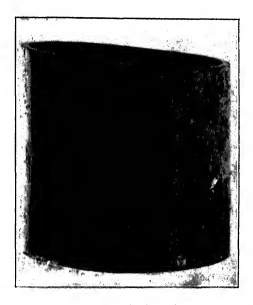


Fig. 22 Roderstube.

paint beneath it. The tubes were pitted with rust in spots and streaks when they were first removed to be cleaned. These show in the photograph, but the corrosion was stopped by the paint. The light-colored scale-deposit was left on part of the tube, and shows on the sides of the figure.

Pieces of iron coated with graphite paint and dipped in muriatic, sulphuric, and nitric acids, and allowed to dry with the acid on them showed at the end of nineteen days no injury to the coatings. Other

paints.

While tests of paints are not regarded by many engineers as indicative of their value to resist the ordinary influences upon a conting exposed to weather, they do show that a conting that can withstand the above severe tests is certain to give more satisfactory results in its general use than the many commercial paints whose low price and not their protective qualities is their principal recommendation. They also show that if the conditions to which a conting is to be subjected are known, it can generally be furnished to successfully meet them.

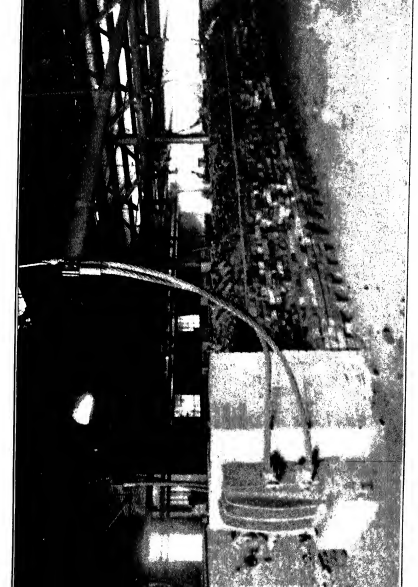
Other tests of graphite paints in competition with other commercial paints are given in the following chapters on paint tests. For a roofing paint the graphites high in carbon, of themselves, or mixed as silien-graphite compound paints, are of marked excellence. They do not harden as rigidly as the iron oxides used for roofing purposes. They endure long exposure to the sun, hence are less liable to crack or flake off, and they follow without injury the expansion changes in the metal they cover. Their darker color and higher cost compared with iron-oxide paints used for roof contings are more than offset by their better protective qualities and longer life.

Electric-furnace Graphite.

However engineers may differ about the respective merits of a low or medium grade of unorphous-graphite mineral for a straight paint compared with a flake-graphite and silica compound, their attention is liable to be attracted in the future to a new product that has entered the field for a pigment, under the name of

"Acheson Graphite."

This substance is an uncorphous-graphite pigment of high-carbon content, whose physical character seems to be materially different from the high-carbon mineral graphites heretofore used for paints. Although amorphous or granular in character, as compared with the other forms of graphite, such as the flake-graphites, it is nevertheless distinctly a graphite product, and contains absolutely no trace of amorphous carbon, the name usually modified to such forms of materials.



Die 23. Aberthe Primase for Matchart att z. Adesen Grayllie.

state of subdivision and of great uniformity in size of the individual particles.

Its amorphous character renders it far less repellent to the oil than the natural graphites containing approximately the same percentage of earlier. This quality gauges it to remain in place in the

Its amorphous character renders it far less repellent to the oil than the natural graphites containing approximately the same percentage of earbon. This quality causes it to remain in place in the oil, and it is not as easily moved out of position by the drying action of the vehicle, as is the case with a high-earbon flake-graphite.

Acheson graphite used with a boiled-oil vehicle will set in the coating without the aid of any inert substance to hold it in place while drying. Used with raw oil, it requires a drier to secure the initial set of the paint, particularly if the coating is to be an external one exposed to the vieissitudes of weather.

Its manufacture is entirely unlike that of any other pigment, and is shown by Fig. 23, illustrating the style of the special electric furnace used to produce it.

In manufacturing graphite in this way, anthracite coal is heated several hours in the electric furnace by means of a powerful electric current, approximating 1000 horse-power of energy. The temperature of the mass of coal is raised to a point where the carbon is converted into carbides of the various constituents of the ash, which in anthracite coal are very evenly distributed. The temperature is then carried to the point where the carbides are decomposed, and the principal constituents of the original ash, silicon, iron, sulphur, aluminum, etc., are driven off as vapors.

The residue removed from the furnace is carbon in the form of graphite, perfectly free from any trace of the amorphous carbon or coal from which it was produced. Its method of manufacture is probably a duplication, upon a small scale, of the process by which the natural graphites were formed in the earth. The purity of the product depends upon the temperature to which it has been raised; for commercial purposes, it contains about 90 per cent of carbon. The 10 per cent of ash still remaining in the carbon is practically as inert as the graphite itself, and intimately associated with it. The furnace product is broken up, and the grades suitable for various purposes

separated. The grade used for a pigment is pulverized to an im-

for an ideal paint.

Unfortunately, there is no standard for a graphite paint as there is for a pure-white or red-lead paint. The consequence is, that where graphite paint is specified by the engineer, he is to an extent working in the dark, and does not feel at all sure but that the coating will be spread from some one or other of the many abominations under the guise of "mixed paint," that has not an atom of graphite of any kind in it. Reputation of the manufacturer or dealer in graphite pigments or paints is quite as essential as in the case of the lead and zine products. Adulterations in a graphite-mixed paint are more easily concealed from the eye than in those having a lead or zine base, and are equally, if not more, annoying to the engineer.

CHAPTER XIV.

BESSEMER PAINT.

A special pigment, claimed to be of the inert class, has lately come into use to replace oxide of iron as a straight paint for ferric structures. During the short period it has been upon the market, its use has been attended with many favorable results. It is a German development, and is reported to be the pulverized slag from Bessemer basic process steel furnaces. It is claimed to be free from the sulphur and phosphoric acid elements that are usually present in iron-oxide pigments.

It is prepared as a mixed paint ready for use. The finely pulverized furnace slag is ground in linseed-oil containing a small amount of one of the copal resins that makes the paint coating very clastic even after long exposure to the sun.

It is claimed to spread easily, covering 1000 square feet per gallon; but to do this the use of short bristle brushes is recommended, the effect of which is to rub out the coating very thin. But however closely the paint may thus be forced into contact with the surface being covered, it cannot be as well protected as where the paint is spread by long bristle brushes, and a sufficient amount of painters' labor and time is given to spread the coating. The pigment is not so deeply embedded in the vehicle, nor so well protected or bonded to the coated surface as it is when spread over a smaller area.

Bessemer paint in its natural color is a very dark gray, though it can be made a lighter shade by the addition of other substances (not of its own nature) to tint it. In this case the coating will be no more durable than the life of the most perishable pigment in the paint, as is the case with all compound paints.

associated pigment or substance from the action of the atmosphere. All slag pigments are electro-negative to the metallic base pigments, and to all of the metals that constitute a part of their composition, also to the metallic surface that is coated with them.

Insulating qualities are claimed for Bessemer paint; but other paints free from metallic oxides also have this quality. The insulating qualities of any paint are due to the vehicle more than to the pigment, with the single exception of india-rubber. In any case, a paint coating cannot resist electrical currents of high potential; to moderate or low potential the insulation would be more or less resistant according to the amount of resinous matter in the vehicle. In this respect Bessemer paint, containing as it does a small amount of fossil resin, would be better than a paint containing none.

Common resin or resin oil should not be substituted for copal; they are not desirable elements in a paint, as they dry hard and

erack the coating or cause it to crumble and rub off after a short exposure in the open air or similable, and they promote corrosion. (See Paint Tests, Chapter XXIX.)

Pulverized mineral wool has been proposed for a pigment. It is a furnace slag riven when in a molten state by a current of steam. But merely pulverizing it impacts no protective value to it for a

a furnace slag riven when in a molten state by a current of steam. But merely pulverizing it imparts no protective value to it for a pigment. It is acid in reaction, electro-negative in character, and when used for covering steam-pipes or other ferric bodies, on becoming damp is a virulent agent for promoting corrosion. A sample of mineral wooi analyzed by Prof. Egleston, of Columbia University, gave the following result:

Substances.	Per cent	in Water	lu Water	
Water	80.1	1.08		
Potash	0.19	0 19		
Soda	1.75	1.75		
Magnesia	19 82	0.12	19.70	
Lime	26,56	1.61	24 95	
Sesquioxide of iron	0.64		0 64	
Alumina	7.84	• • • •	7 84	
Silien,	38.97		38 97	
Sulphur (mostly as a sulphide,				
probably, of calcium)	2.64	0.32	2 22	

Buluble

Insulithia

analysis of blast-furnace slag, and with it may be taken as representative of this class of substances.

No analysis of the Bessemer pigment is given by the manufacturers of the paint. It is, however, supposed to be a tetra-basic phosphate of lime, containing about 20 per cent of phosphoric acid and 50 per cent of lime, associated with other mineral and metallic substances in Bessemer iron ores. Some of these substances are partially consumed in the working of the furnace, and the balance fluxed off as Bessemer basic slag.

Bessemer basic process steel was made by the Pottstown Iron Co., of Pottstown, Pa., for a few years previous to 1893, and the pulverized slag was sold for a fertilizer. Since 1893 the basic process for making steel has been suspended in America, and the slag is now procured from Germany.

ANALYSIS OF BESSEMER CONVERTER BASIC SLAG.

Phosphoric acid	21,37	per	cent.							
Lime,	45,26	·u	**							
Iron exide	12.00	"	££	{	Equal meta	to Illio	8.40 iron.	per	cent	of
Silica	5.10	**	"	•						
Magnesia	5.90	"	u							
Alumina	4.01	**	u							
Soda and potash	0.80	"	u							
Manganese oxide	5.56	u	и							
1	00.00	**	4.							
Assumption of Many Co.	*** * * * * * * * *	1	, w 4	٠١.	. *****	n ,	e on	1111		J.,,,

An analysis of blast-furnace slags, the mean of 2000 samples from furnaces working on gray forge pig iron, is given for a comparison with the Bessemer converter slag.

with the Bessemer converter	Blug.			
Silien	43.07	per	cent.	
lime,				
Alumina	14.83	"	u	
Iron oxide	2.83	u	"	Equal to 1.98 per cent of metallic iron.
Peroxide of magnanese	1.37	"	et	•
Mngnesia	5.46	"	u	
Potash	1.84	et	et	
Calcium				
Sulphur				

Blast-furnace slags are acid in reaction, while Bessemer slag is basic or neutral. Both are pyrogenic bodies unaffected by heat or sunlight, and neither is oxidized by the atmosphere.

A German chemist* gives the analysis of Bessemer paint, as known to the trade in Germany, as follows: "The pigment contains baryta, alumina, iron oxide, lime, silica, zine oxide, sulphuric neid, carbon dioxide, and phosphoric acid." No definite percentages of these substances are given in the analysis, nor any mention whether they were separate constituents of the paint assembled in the process of grinding and mixing, or that any number of them were found combined together as a single pigment. "Graphite or other earbon is used as coloring matter, and linseed varnish as the vehicle, turpentine constituting the drier. The presumptive constitution is, therefore, lithopone, or silicious calamine ore, containing baryta and chalk, together with graphite or other form of carbon, and linseed varnish (with probably turpentine as a drier). When treated with hydrochloric acid it disengages sulphureted hydrogen."

There are many formulæ for compound pigment paints in this country, each of some declared excellence by the manufacturers of them, if not by the users; but it is difficult to select one that for the varied composition will equal this German product. Whatever may be the composition of the American brand of "Bessemer paint," it appears, from the above description, to be wholly unlike that of its German namesake, and is certainly superior to it for a ferric paint.

^{* &}quot;Andes' Iron Corrosion."

CHAPTER XV.

NATURAL-ROCK HYDRAULIC CEMENT.

Hydraume-cement coatings, either in the form of a plaster coat laid on by a trowel or as a wash or brush coating, have not been much used for the protection of ferric structures as a substitute for paints, though its use as a protection from corrosion of iron embedded in masonry is common and its value for this purpose under certain conditions is recognized.

Hydraulic cement made from the ground mineral varies greatly in quality, its general composition after calcination, that makes it caustic and anhydrous, being:

Silica, sand, magnesia, sulphur, and many metallic oxides are also present in some amounts in many varieties of the hydraulic mineral, all of which affect the quality of the cement unfavorably when it is used for a mortar, and are more objectionable when it is to be used for a coating on ferric bodies.

The adulteration of mineral hydraulic cement is generally, from the same class of uninerals of inferior quality, with free sand, silica, and iron ores containing sulphur in the form of sulphides, and all are imperfectly roasted and pulverized.

Their setting quality and strength are very irregular and uncertain whatever their trade name, or the manufacturer's report of the large quantities sold

Alumina (Al_2O_3)	7.26	11	44
Magnesia (Mg.O)	3.10	11	it.
Iron oxide $(\lceil c_2()_a \rceil, \ldots)$	3 24	11	66
Insoluble residue(clay and sand)	0.11	1.6	4.6
Sulphur anhydride(SO ₂)	1,36	ii	(i
Carbonic anhydride (CO ₂)	0.33	"	44
Water (H_2O)	1.97		46
Soda $(Nn_2(1))$ and loss Potash $(K_2(1))$	0.04	14	16
•	100,00	44	11
Tensile strength of the neat cement at the	end of	7	days equals

613 pounds; at the end of 28 days, 800 pounds, with one part of cement and three parts of sand for the same periods, 228 and 360 pounds.

The composition of a natural-rock hydraulic cement from Chatta-nooga, Tenn., is

Silica	22.17	per	rent.
Lime	65.68	* *	
Alumina	8,20	* *	14
Magnesia	1.45	**	11
Oxide of iron			
	100.00	11	**

It is a natural Portland cement similar in character, but superior to that found at Boulogne, France. The Chattanooga deposit is in the form of layers, and is over 50 feet thick. After calcination at a white heat, the following are the average results from a number of tests of briquettes.

After an exposure of one hour in the air and 23 hours in water, the tensile strength averaged 235 pounds. After one day in the air and 6 days in water, 623 pounds. After one day in the air and 27 days in water, 797 pounds. It is the strongest natural-rock cement in the world.

Portland cements are commonly made from a dual combination

or slag sand.

These substances are pulverized and mixed in some proportions that vary with the different manufacturers. The mixture is then calcined to nearly the point of fusion, or to actual fusion, forming a cinder which is finely pulverized and called Portland cement.

Furnace slag, the waste product from blast-furnaces (see Analysis, Chapter XVI), is also used as the base of Portland cement.

The slag is heated in mass and quenched in water to granulate it, making slag sand; then dried and mixed with about one-fourth part of slacked lime, and hen finely pulverized to form the cement. Slag sand usually contains from 0.5 to 1.5 per cent of the sulphide of iron, that has a tendency to oxidize on exposure to the air, which action is destructive to cement in above-ground situations. Slag cements are from 6 to 8 hours in setting, as against 1 to 3 hours for the American brands of mineral cements under test of a one-pound needle.

The color of slag cement is a delicate lilac or almost white. If the slag sand has been roasted prior to pulverizing, it is generally of a dark color similar to the ordinary brands of dark Portland cement. If a greenish color is present in the cement, it denotes the presence of a large quantity of the sulphide of iron. This greenish color is also found in some brands of the ordinary Portland cement, where the substances from which it is made contain iron sulphide, and when there has been a deficiency of heat in the oxidizing flame of the kiln.

Slag cements, when mixed and exposed to the air, must be well covered, else they will crack, though they harden under water without swelling or any material change in volume. They are completely hydrated during the process of manufacture, do not require aging, and do not deteriorate in storage.

The character of furnace slag and all the processes of its manufacture into cement require as close attention to secure a reliable product as is required for the mineral or Portland cements. The engineer should select a cement for a wash coating for ferric surfaces, or for a mortar, by its properties, not its name, and should require the standard he desires to reach to be met by systematic and rigid tests of every invoice of the cement, and many tests from each in-

The nature of the cement has much to do with its effectiveness. The quick-setting cements require the most care in their application, and are generally the best for ship work. If the thin cement wash once sets in the bucket it will not again set if stirred up it is then useless. Constant stirring of the paste is necessary, as fifteen to twenty minutes after mixing suffices for it to set if not kept constantly stirred.

The Portland coments set slower than the American or natural mineral cements. Quickline is sometimes added to delay the setting, but renders the cement more caustic and destroys the protective qualities of the vehicle in the underlying paint, and opens the way for moisture in the cement to reach the metal, and all of the contings soon peel or flake off either from the corrosion of the iron or by the destruction of the bond between it and the paint.

Coment coatings, unless spread where they are freely exposed to a circulation of the air, are damp, and being porous, are not proof against the penetration of gases or liquids. If by accident they are exposed to the action of any copper scales, scrap-metal, or water charged with acid or alkaline substances, the soluble salts of copper thus formed will penetrate the coating, deposit the copper upon the iron or steel surfaces, set up a galvanic action, and corrode the metals beneath the coment coating.

The hardness and rigidity of cement contings render them liable to flake off the metallic surfaces under strains due to changes in temperature of the metal that the cement cannot follow. Such places, though of minor extent, are generally inaccessible, and are quickly corroded; this action being hastened by the difference in potential between the exposed metal and the cement coating, even under ordinary atmospheric conditions. It is more rapidly developed if acidulated solutions or vapors are present, as they nearly always are abourd ships.

All of these disadvantages in the use of cement can with proper care be in a great measure lessened, if not altogether avoided. Cement coatings are in many cases the only protection that can be used to prevent corrosion, or to arrest it, even where it has progressed to an extreme or dangerous point.

The collapse of many stand-pipes shows in nearly every case that corrosion was the principal cause of their failure, as its progress at one or more places had reached such a degree that it only needed a small extraneous disturbance to wreck the structure. A case in point is that of the 60-inch-diameter wrought-iron inlet-pipes 300 feet long, and the lower sections of the stand-pipe of a large waterworks erected in 1860 and in use only a few years, when corrosion had developed upon the inner surfaces of the pipes in so many large spots and blisters, in such an irregular manner, that the engineer reported "that over one-half of the strength of the pipes to resist external pressure had been destroyed. Parts of the pipes were un-affected, the mill-scale and shop-marks being in place, while nearly

The inlet-pipes being buried in river-silt containing a large amount of clay, were comparatively unaffected, though below the water-level; but it was still water, not subject to motion like the suction and force sides of the pipes.

one-half of them presented an appearance of being inoculated with

poison."

The inside surfaces were scraped clean as possible, and then coated with one coat of neat hydraulic cement from ‡ to ‡ inch thick, laid on by a trowel by house-plasterers.

The water-tower was wrecked by a tornado in 1800, and all the pipes were found free from rust in any degree, and probably would have lasted indefinitely.

Imported Portland cement was used on one part of the pipes and Louisville cement for the rest. The former set slowly and had an indifferent adhesion to the iron. The Louisville cement set promptly and was easier to apply.

Many other instances of the successful use of hydraulic cement in similar situations could be cited. The quality of the cement, the manner of mixing and applying it, and the personal equation of both the engineer and the employé, are factors for success. Failures of cement coatings are more frequent than successes for the reason that in their application one or more of these requirements have been neglected.* Prof. J. M. Porter of Lafayette College divided a sam-

be made of it in a mortar, one part cement to three of sand, according to the rules recommended by the committee of the American Society of Givil Engineers. The resulting average strengths reported from each of the nine samples were as follows: 75, 102, 114, 136, 153, 163, 176, 225, 247 pounds tensile strength per square inch. Average for all the samples was 153 pounds, and the lowest strength was but 30 per cent of the highest.

If these results produced by experienced men in permanent laboratories vary so much with the same cement, what is to be expected from the inexperienced and careless laborers who are generally employed to mix and apply concrete, mortar, or cement coatings?

Neat hydraulic-cement contings crack, they set so rapidly that there is always a probability of their setting before the workmen can spread them, and the tendency of the workmen to "knock them up" when they indicate setting or have set, instead of mixing a fresh batch, is almost irresistible, the result being a conting of very uncertain character,—streaks of firm and close-clinging cement alternating with those of dead cement, that readily yield to a slight change in the temperature of the metal or covered surface, or to a slight mechanical injury or a frost. A strong heat from the sun also causes them to flake off.

Bad milling and insufficient burning are a frequent cause of poor cement, also an excess of magnesia in the limestone or added adulterant. Magnesia causes a chemical change or disintegrating action in the cement when wetted in the unixing. Free, natural sulphate of lime is a dangerous impurity. A low specific gravity and a dark-brown color are indicative of poor burning, and are easily detected.

For the protection of iron or steel beams or grillage, laid as the foundation or structure work below the water-line to be embedded in cement concrete, the metal should be bright and free from nill scale, which is an electro-negative element, and with the moisture present, is certain to produce a galvanic couple with the iron it covers and promptly start the corrosion, that will proceed uninterruptedly so long as any metal is left for it to act upon. Every atom of the red rust as it forms, being also electro-negative, increases the galvanic

energy on the remaining metal. The rate of this corrosion will probably reduce the beams in less than one hundred years to a condition where their strength to sustain any incumbent load will be no greater than an equal quantity of tan-bark.

No paint coating on the metal can resist the galvanic action, and it should be applied solely to prevent any slight corrosion that may occur from the time the metal is cleaned until it is laid in situ.

A lampblack and oil, or a graphite paint are good anti-corrosive coverings, but best of all coatings for these situations is a refined bitumen and dead-oil mixture applied hot. Oxide of iron, or any other coating containing electro-negative substances that induce corrosion under atmospheric conditions, will only add to the strength of the galvanic couple, by bringing their oxidizable elements into the field. The nature of the soil in which the metal is directly in contact will also contribute to the corrosive action through the galvanic couple, blue clay or solid rock only excepted.

Concrete, as generally laid, is very porous, and is seldom so proportioned, mixed, or rammed in place as to enable it to fill the voids in its mass, and capillary action will enable the moisture and soil acids to reach the metal. In all such foundation work the cement should be of the best quality, free from sulphur elements, the filler should be of small size, and the sand absolutely free from salt or sea-sand in order to minimize the dangers of corrosion. These precautions are seldom if ever taken, even in part, much less as a whole. The neglect of these particulars will soon be apparent in the failure of many an important structure whose life will be measured by a few decades instead of centuries. That the corrosion in these cases is out of sight and mind and irreparable will be the more aggravating.

Porosity of Hydraulic Cement,

In a general way, engineers and architects are inclined to blindly trust hydraulic cement in many locations where in parallel cases it has failed. The quality of a cement suitable for a concrete block would not be advisable in a wash coating for a wall or to bed an auchor

by ashlar masonry, were disintegrated in iess tigor a year by the action of sea-water. There was a movement or change in the character of the beton, even when 660 to 880 pounds of Portland cement per cubic meter of sand was used. In parts of the work where they had not been exposed to the action of sen-water, the beton of all proportions of cement and sand were not ordy very porous but very permeable. Under a head of about three feet, the permeability was indicated by a rapid fall of the water in the vessel where the beton block was under test. The permeability of the cement was in all cases accompanied by a disintegrating effect in the beton. The disintegration was found to be due to the formation of perceptible quantities of the sulphate of lime by the action of the sea-water on the Portland cement. The sulphate of lime, when formed in the mass of concrete, solidified more or less completely in crystals of such a unture us to develop considerable molecular activity. Some of the beton cement analyzed .75 to .80 per cent of sulfduric acid. The greater the amount of water used to mix the cement the greater

The greater the amount of water used to not the cement the greater was the permeability and porosity of the concrete, even with all proportions of the cement and sand. Mortar made with 7 per cent of water was very permeable, and this increased perceptibly as the percentage was increased to ten and eleven. In all cases where secwater instead of fresh water was used to gauge the cement, the land effects in the mortar were at once apparent.

Prof. Bauschinger's experiments showed that the adherence of a first-class Portland cement to a bright wrought-iron floor beam was 625 pounds per square inch; that mixtures of two parts of fine, sharp bank sand to one of cement reduced the adhesion to about 70 per cent of the above value. In mixtures of three parts of sand to one of cement the adhesion value was less than 50 per cent. That with each increase in the percentage of sand from the above amounts, the reduction in strength and adhesion was very rapid. The quality of the cement had a great effect upon the adhesion value. In the commercial cements usually provided for contract concrete, the adhesion was frequently only 20 per cent of that given above.

Bloxaur's "Chemistry," edition 1895, 190, 376, 377, states "that

the ordinary corrosion of iron is accomplished only in the presence of

absorbs oxygen from the atmosphere, in accordance with

$$2 \text{FeCO}_3 + O - \text{Fe}_2 O_3 + 2 \text{CO}_2, \dots, \dots, \dots$$
 (i)

The Fe_2O_3 combines with the moisture and is deposited as $2\text{Fe}_2\text{O}_3.3H_2\text{O}$, or ordinary iron rust. Iron in its ordinary state is not affected in perfectly dry air, and it will not rust in water containing a free alkali or alkaline earth or an alkaline carbonate, because the affinity of these alkaline substances for any acid is greater than that of iron, so that they would neutralize the acid before it had time to attack the iron."

This neutralizing action, however, would only be effective for a short time or until the alkaline substance became saturated with the acid element. There are no locations where concrete is used or cement contings applied to iron for its protection from corrosion where reviving the saturated alkaline substance is possible. It is therefore only necessary to have a limited amount of some acid present with air and moisture to cause the ultimate destruction of a large amount of iron, because the CO₂ or other acids present never become fixed, but are always active, passing from molecule to molecule, as long as there is any free metal for them to attack.

It is proposed to increase the safeguard afforded by alkaline substances to delay corrosion by mixing the concrete, mortar, or wash coating with whitewash instead of plain water. The small amount of lime thus added to the cement does not materially detract from its strength.

Slag cements, because of the sulphides present, should be avoided for use in concrete or any coatings in contact with iron. It is hardly possible to assemble them with an amount of any alkaline substance that will permanently neutralize the acid element present in their composition, aggravated in nearly every instance by the porous nature of all concrete constructions caused by deficient ramming to fill the voids occupied by the enclosed air, also by the surplus of water used in mixing.

Even a cement free from the sulphur element, if mixed with a small quantity of einder, or if laid in soil containing einders or pyrites, will absorb the acid and collect it in dangerous amounts in the voids of the government. Once there it will ultimately reach the metal and

further aggravated by the fact that nearly all ground-water is charged to some extent with saline or sulphur elements or both, that would soon saturate any alkaline substance present in the cement. When this point is reached corrosion of the grillage will inevitably ensue even if the imposed columns show no evidence of its action.

Grillage ironwork has been removed from concrete foundations laid only five years and found to be corroded & inch or more over its whole surface. The thickness of grillage beams is seldom & inch, so that thirty or fifty years will practically limit the sufety of many of the modern steel skeleton structures.

The protection afforded to steel by Portland cement has been subjected to experiment by Prof. Charles L. Norton.* "Two brands of American cement were selected, tested chemically and physically and found to be good. A sharp, clean bank sand and fragments of traprock and flint were thoroughly washed and used for the concrete. The cinders were washed and dried; they tested distinctly alkaline with a small amount of sulphur. All the ingredients were mixed dry in every case, and when wet with a minimum amount of water were tamped until they flushed.

"Briquettes were made in duplicate with both cements, viz., neat cement, one part to three of sand, one part to five of broken stone; cement one part, two of sand, and five of stone; cement one part, sand two parts, and five of cinders. Specimens of mild-steel rods 6" + 4" diameter, mild sheet-steel plates 6" + 1" + \frac{1}{32}" thick, and strips of expanded metal 6" + 1" were all cleaned bright. All three pieces were put into each briquette and were enclosed in separate tin boxes, which also contained a specimen of each metal unprotected. One half of the briquettes were set in water for one day and the rest for seven days before scaling them up tight. One-quarter of the boxes were then subjected to each of the following exposures. To an atmosphere of steam, air and carbon dioxide; to air and steam; to air and carbon dioxide, and the other samples set upon a table in a room with no special care as to their temperature or dryness.

"At the end of three weeks the briquettes were cut open and the

The location of the rust spot was *invariably coincident* with either a void in the concrete or a badly rusted einder. Rust had as usual produced rust.

"In the more porous mixtures the steel was spotted with alternate bright and rusty areas, each clearly defined. In both the solid and porous cinder concrete many rust spots were found, except where the concrete had been mixed very wet, in which case the watery cement had coated nearly the whole of the steel like a paint and protected it.

"Some briquettes made of finely ground einders and cement in varying proportions up to one of cement to ten of cinders and exposed to moisture and carbonic neid showed how effectually the presence of the cement prevented rusting, provided there were no cracks or crevices or distinct voids. The corrosion found in einder cement appeared to be mainly due to the iron oxide in the cinders and not to the sulphur. Cinder concrete, well rammed when wet to fill the voids, is about as effective as stone concrete in protecting steel."

These latter conclusions would depend greatly upon the absence or low percentage of iron oxide and sulphur in the cinder. To render these elements inert to iron, there must be enough free alkaline substance in the cement to saturate the acids without disturbing the general composition of the cement as a binding element.

If the metal is painted before the application of the concrete, whatever its composition, the continuous void left over the whole surface of the metal by the decay of the paint is the best possible condition for inaugurating corrosion. Air and moisture will find ready access to this void, also to the voids left by building bricks and terra-cotta blocks, the porous nature of which are favorable to the condensation and absorption of moisture and atmospheric gases, that are more highly charged with corrosive elements in cities, tunnels, subways, and other locations where the use of structural steel work is in most demand.

How far the protection of ferric foundations, either near or below the water-line in the many structures already built, or in progress in all parts of the world, has been considered by their architects and engineers time only will reveal. For those proposed, like the miles protection from corrosion must be employed than has ever been adopted. No wash or trowel coating of cement, good or bad, or applied in mass, will avail for but a short period to protect the metal that these structures must rely upon for agreat part of their strength, The hardness and inelastic character of cement or mortar coatings

will cause them to crack under the vibrations inevitable to all railway structures; and while resisting water in mass, they will absorb moisture sufficient to be always damp and in that condition are of the least strength. The wires of the anchorage ends of the cables of the Ningara Falls

suspension bridge were opened for a short distance where they entered

the anchorage pits. These ends were embedded in hydraulic cement. and at the end of forty years many of them had become so corrolled that the strength of the structure was seriously impaired. The corroded strands were replaced by new wires, and the top part of the anchorages opened to allow the cement work to dry out and remain In this case and with all ferric material embedded in concrete, the caustic action of the usual make of coment, whether damp or wet, will furnish the earbonic acid necessary to destroy any linearly oil coating or paint that rovers them and induce corresion. The subsequent drying out of the cement coverings only delays for a short time the ultimate destruction of the metal.

Iron anchor burs and chains embedded in concrete below the waterline for 100 and 200 years were free from rost when removed. A reshwater immersion, no access of air, no acid element nor iron exide or calcium sulphate, also no voids in the cement was the secret of their perfect condition.

The present method of constructing buildings wholly or in part of steel framing and concrete, avoiding the use of brick and stone nursonry as far as possible, is causing a great deal of anxiety among architeets and engineers as to the future state of the metal so embedded. That metal needs some additional protection from the caustic action of the impure cements too frequently employed, also from the quicklime mortar, beyond the usual cont of paint, is recognized. At a late meeting of the English Architectural Association, Mr.

H. Humphrey gave as the result of his experience that metal buried

other compound containing pitch and sand; that some samples of cinder concrete analyzed as high as three-fourths of one per cent of sulphuric acid. A case was cited by another member of the association where a hot-water pipe laid in cinder concrete was rotted away in a very short time.

The einder concrete used in the floors of the steel-frame sky-scrapers in New York City invariably shows the presence of sulpharic acid strong enough to redden litmus-paper.

Gas-pipes embedded in plaster of Paris (gypsum) have been found to be completely rorroded in a few years. The use of gypsum in comput to hasten its setting is detrimental. Gypsum is soluble to some extent in water, besides it contains water from its hydration, which absorbs carbonic acid from the air that quickly rauses corresion. The rust so formed absorbs moisture and carbonic acid and further hastens the corrosion.

The screwed ends of all pipes are invariably attacked. They are of bright metal only about $\frac{1}{16}$ inch thick and seldom, if ever, have even a brush conting of any paint to protect them when put up or left in place. Galvanizing the fittings and body of the pipes does not protect the screwed ends; the corrosion at these points is only hastened by the galvanizing.

The effect of corrosion upon the floor beams and other structural parts used in modern architectural work has been the subject of discussion by the American Society of Architects, the concensus of their opinion being expressed by one of the prominent members as follows:

"With regard to the strength of the steel-enge constructions, both as to wind strain and other disturbing strains, there is no question. All objections arising from these points have been overcome, but unless exceptional care is taken in the construction to protect the steel cage, particularly at its joints, from corrosion, this class of buildings will not be permanently safe. It is perfectly feasible, with great care, to protect the steel frames from corrosion, but I am convinced that many high buildings have been put up in this country where the proper cave in this respect has not been taken nor the necessary

stories, the paint was dead and corrosion established. This was particularly noticeable in many portions of the beams where the usual top dressing of coal cinders bud been hid to level up the arches forming the foundation for the artificial stone sidewalk. The rivets that held the corner angle-irons to the beams were nearly all loose from the corrosion around their heads or points and had lost their set or draw.

In marine work, hydraulic cement is used almost exclusively as a brush coating on the inside surfaces of the ship's frames and plating in the lower holds of the vessel where the metal is exposed to the action of bilge-water, alkaline and acid solutions from acids, and leakage from the eargo liquids. One or more wash coatings of cement are applied over the red-lead, black-varnish, or other oil paint coating laid on during the construction of the ship, and that generally serve to protect the metal during this period. The coatings in the lower part of a ship are damp by reason of the confined saturated sea air, but the cement (if good) forms a close, clinging coating that seldom fails unless by mechanical injury or improper mixing or application, and is easily repaired if injured.

The confined spaces abourd a ship almost preclude the use of an oil paint or varnish, however quick drying it may be, without the use of forced ventilation to provide the oxygen necessary in the drying of paint. Such ventilation is practically impossible in a ship at sea, or in most cases in dry dock.

Cement for rendering, with the object of making brickwork watertight, should be mixed with an equal part of absolutely clean sand free from salt or sea-sand. Cement for any use should be carefully turned over by the shovel and exposed to the air before being mixed or wetted.

Brickwork is one of the worst surfaces to hold a paint, good or poor. A hard-burnt brick will absorb 8 ounces of water, a salmon brick nearly 11 ounces. Brickwork absorbs toost of the oil in the paint, leaving the pigment on the surface of the bricks without sufficient bond to hold it, and it peels in strips. This peeling is hastened by the caustic action of the cement or lime mortar and the soluble salts in the sand of the mortar, which soon destroys the organic mat-

- crustaccum, is represented by Fig. 24.*

Walls a year or more old are less troubled with the efflorescence or by the peeling of the paint from the fungus.

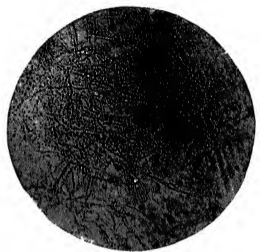


Fig. 24.—Photomicrograph > 600 of *Penicilium crustaceum*.—This is the greenish fungus which makes calcium oxalate on brick walls.—(M. Toch.)

When walls are freshly laid or plastered the surfaces can be prepared for painting by applying a solution made of twelve fluid ounces of sulphuric acid in a gallon of water and repeating the application when the first one appears dry. Allow the coatings to stand for a day or two, then riuse off with clear water, and when dry prime and paint as usual. This process changes the lime in the mortar and cement from a caustic carbonate to a neutral sulphate of lime; also produces a uniformly absorbent surface free from spots that are more porous than the general surface, or that contain lumps of improperly slaked or mixed lime. The surface so prepared takes the paint easily and well and does not blister nor peel.

If plastered surfaces a few months old be washed with a solution

^{*} Maximilian Toch (New York City). Journal of Chemical Industry (London), Vol. XXI, No. 2, Jan. 31, 1902.

peeling.

A silicate of soda solution made from equal weights of silicate and

A sincate of soon soutton made from equal weights of sincate and warm water, and applied with a brush, is also recommended for preventing the pecling of paint on walls, but for outside exposures it is not so effective as the above acid treatment.

Waterproofing Bricks and Sandstone. At a recent meeting of the Australian Association for the Advancement of Science, Professor Liversidge read a paper on the "Waterproofing of Brick and Sandstone with Oils." Experiments were made with the view of ascertaining the length of time that brick and sandstone are rendered waterproof or protected by oil. The oils used were the three commonest and most readily obtainable for such purposes, viz., linseed oil, boiled lipseed, and the crude mineral oil known as "blue oil," used for preserving timber. The weatherings were made upon a flat portion of the laboratory roof fairly exposed to the sun and weather. Good, sound, machine-made bricks were experimented on. The amount of oil and water taken up by the sandstone was very much less than that absorbed by the brick, although the area of the sandstone cubes was much greater than that exposed by the bricks. Equal amounts of raw and boiled oils were absorbed; the blue oil, however, was taken up in much greater quantity by both brick and sandstone, but by the end of twelve months the whole of the 134 owners of blue oil had apparently evaporated and the brick had returned to its original weight. The bricks treated with raw and boiled oils remain unchanged. After the second oiling in November, 1890, and exposure for nearly four years and two months, they had practically retained all their oil. inasmuch as they had not lost weight, and were also nearly impervious to water. It was noticeable that the sandstone cubes treated with raw and boiled oils returned to their original weights, but did not appear to have lost the beneficial effects of the oils, being also practically waterproof.

Portland or other hydraulic coments free from the sulphate of lime, when mixed with about 15 to 25 per cent of a red-lead paint, forms a tough elastic coating that dries hard enough to resist the action of locomotive exhaust steam and cinders on the surfaces of iron leanns, trusses, and the buckle-plates of low headway bridges. It is also

favorable situations. From a government analysis of it, the composition is approximately 30 per cent of an oil vehicle, 65 per cent of refined special bitumen and selected fossil resins, and 5 per cent of a carbon pigment. It is laid on or spread like a thin coating of mortar on brick masoury or plastered walls. It adheres firmly, becomes very tacky, and can be plastered over with cement or lime-mortar coatings that adhere firmly. When these plastered coatings are dry they can receive an oil paint of any desirable color, unaffected by dampness from the walls.

A grade of the "R. I. W." is also made to apply to damp walls not intended to be plastered, also to iron structural work. This is applied with a brush, and contains more pigment than the trowel grade. It is thoroughly damp-proof, and receives oil-paint coatings without any tendency to craze them or to peel.

A grade of this composition, to be spread with a brush on the inside of tanks where acid and alkuline solutions are stored, effectually resists the action of these liquids. In chemical works for the protection of the ironwork and other metals, it has shown great resisting power. A special instance of the waterproofing character of this compound to resist the action of running water under a considerable head is on the concrete monolithic water-power house on the St. Lawrence River at Musseuu, N. Y., where several thousand square yards each of the trowel and brush contings were applied, and completely corrected the porosity and permeability of the cement walls that were seriously endangering the structure.

Herr Wm. Cremer, superintendent of the gas-works at Enskirchen, Germany, states: "That the unimoniaeal liquor from gas-works, even in the weakest solutions, detrimentally affects the cement and tankwalls exposed to its action. Coating the surfaces thus exposed with liquid glass (tangstate of soda) protects the cement, also renders the surfaces quite leakage-proof even in very old work."

CHAPTER XVI.

BOWER-BARFF COATINGS.

"Bower-Barff" is the name given to the rustless coating formed upon east iron, wrought iron, and steel, when exposed to a low red heat in special ovens, furnaces, or retorts, and subjected to the action of superheated steam, carbonic-oxide gas from coal-fires or gas producers, hydrocarbon and hydrogen gas alternately or in combination, according to the several processes invented by Bower Barff, Wells, Gesner, and other inventors.

The original inventor of the rustless iron conting was Prof. Frederick S. Barff, of Kilburn, England, who published an account of his process in 1876, and read a paper describing it before the Society of Arts, London; but the process did not prove commercially successful on account of its high cost and the difficulty of obtaining uniformity in results.

Messrs, George and Anthony Bower, of St. Neats, England, insproved the process of Prof. Burff, and patented it. The right to use it in the United States was acquired by Mr. George W. Maynard, of New York. The first furnace was creeted at the Heela Architectural Iron Works, in Brooklyn, N. Y.

The next and most important of the improvements in this process was invented and patented in 1888 by Mr. W. T. Wells, of Little berry, N. J., who discovered that red-hot iron, in the presence of mingled steam and carbonic oxide, would form the magnetic or black exide (rustless coating) of iron, Fe₃O₄, without the intermediate formation of the sesquioxide, Fe₃O₂ (red rust), the reactions being. 31 e : 411,O, = Fe₃O₄+4H₂. This process is the foundation for all subsequent improvements of the process and is applicable to all forms of east, malleable, wrought iron, and steel where the surfaces are not to be subjected to hard friction or wear, such as bending, hammering, chipping, or other rough usage.

The speciment of the state of t

lation that would injure the continuity of the coating must necessarily destroy the coating. Wherever the coating is broken the metal will rust, though the rust will be localized, and will be greater than the same exposure of the metal not coated, owing to the difference in potential between the two surfaces.

These rust-spots seldom spread or raise the adjacent coating, as is commonly the case with paint, or enamelled coatings. All drilling. fitting, serew-cutting, etc., of the metal should be done before it is put into the converting-oven. In riveting, the oxide in the immediate neighborhood of the rivets will be broken, and bolting together of parts to be connected together must be substituted. In work that is riveted up before being coated, the set or draw of the rivets will be released by the heat of the furnace. This, in the case of light grill, lattice, or fence work, is possibly of small moment, but in work subject to the action of liquids or gases it cannot be ignored and other methods of joining the pieces must be adopted. Shearing, flanging, sharp bending, or driving of unils through sheet-iron roofing, necessarily exposes the metal, and local corrosion of the injured part follows. The bite of the vise or pipe-wrench in fitting rustless serewed steams or water-pipes injures the coating unless special cure and tools are used to prevent the injury. The screwed ends of pipes and fittings are injured if the joints are made up dry, but with red lead, graphite, or other good pipe-joint cements as hibricants, they seldom give trouble if moderate care is exercised in the work.

In cast-iron pipe with bell and spigot joints, the lead packing can be calked without injury to the "rustless" coating by using the round-nose calking-tool, instead of the usual sharp-edged tool that chips the coating. Rustless pipe coatings do not appear to draw in the lead joint any more than the usual coal-tar-dip coatings, from the changes in temperature that all pipes are subjected to when buried in the ground.

The mechanical finish of the metal to be coated determines to a great extent the mode of treatment. Articles in the rough, from which the skin has not been removed, require a longer exposure, higher heat, and a more energetic oxidation than those whose sur-

handsome appearance of the oxidized wate is desired, the suttaces must be cleaned by the sand-blast, or by pickling, and the same care used to remove all traces of the pickling acid by a warm lime water bath and repeated washing with cold water-jets under pressure, as in the case of cleaning the metal for painting (Chapter XXVIII), Foundry-sand upon eastings, if not removed, bakes in the furnace to a reddish-brown color, producing unsightly spots, but does not impair the rustless character of the coating, and unless the coating is to serve as a finish, without being painted, the spots are of no moment; otherwise the sand must be removed to the clean scale surface before treatment. All blow-holes and other defects in eastings must be plugged with brass or iron plugs. Lead or other soft fillings are detrimental to the action of the furnace in producing a reliable or fine-appearing coating, which should be a pleasing bluegray or blue-black color. If the metal is polished before treatment, it acquires a lustrous chony-black finish, very desirable upon certain kinds of articles.

The iron or steel articles treated, owing to the annealing action while in the furnace, are permanently expanded about ${}_{2}{}^{4}_{4}$ inch per foot, for which allowance must be made where this addition will be repeated, as in stair-stringers, columns, etc.

The limit of elasticity of the oxide conting is practically the same as that of the metal it covers. The conting adheres firmly under tensile, torsion and compressive strains, until the elastic limit has been reached, and no further.

In Sir Joseph Whitworth's tests of specimens of Bower-Barffed wrought iron, submitted to tensile strain, small pieces of the oxide coating scaled off when the strain reached 28,618 pounds per square inch, or beyond the elastic limit, and about one-half of the ultimate strength of the specimen. In the case of cast iron, the coating remained in place uninjured when strained to the point of rupture.

Bower-Barffed articles can be heated to temperatures approximately 400° Fahr, and then immersed in cold water without injury. They resist the action of sea air, sea water, sulphurous, and other gases, ammonia, and all alkaline and organic acids in moderate solution, also the caustic action of line and hydraulic capacit, either dry

protective contings under different exposures resulted, viz.:*

CHANGE IN WEIGHT OF WROUGHT AND CAST IRON WITH DIFFERENT PROPECTIVE COATINGS AND UNDER DIFFERENT CONDITIONS, IN POUNDS PER SQUARE FOOT OF SURFACE PER ANNUM.

Whought-from Sheets (No. 23 Gauge, Black).

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Bower-Barffed, Timed, Nickel-plated, Galvanized, Barffed, Black o.c., unprotected, Copper-plated.	gain, .002.0 .0 gain, .000, l	gain, .000,3 '' .1000,1 '' .000,5 gain, .001,1 '' .022,6 '' .005,0	.006.7 .019.4 .050.4 .045.9 .083.9 .137.0 .179.0	.003.6 .007.1 .003.1 .080.5 .117.0 .109.0 .182.0	.002.5 .006.2 .013.5 .042.0 .051.2 .082.5 .091.6
Average gain	.000,2	.005.1	.074.0	.080.3	.040
manufacture of the second of t		Sporterior 1, 200 - 1983	eg. Lyc Tourse out the filter for the same and the same of the	THE PARTY NAMED IN	to in this have a surriging recepting

CART-HON PLATES.

Protective Coatings.		the weather	Immerse	Average	
t tresser i i i vegeregge.	Canada.	New York State.	Fresh water.	Sowage.	galn.
Bower-Barffed. And paraffined, Galvanised, Tinned. Nickel-plated Copper plated Black re, unprotected	0.100, miny 0.000, 0.00, 0.140, miny 0.140, miny 2.1400, miny	gain, 1803.1 .001.0 .001.0 gain, .003.1 	gain, 105,5 " .000,2 " .049,1 " .065,5 " .131,7 " .150,8 " .148,3	.001,4 .008,4 .001,0 .061,0 .083,3 .119,2 .272,4	gain, .002.8 " .002.8 " .027.5 " .011.1 " .053.5 " .067.8 " .106.6
Average gain	.102.9	.002,1	.007.2	.080.7	.0-11

The cost of applying the process must necessarily vary with the size, weight, and other characteristics of the article to be treated. For builders' hardware and that class of articles called shelf goods, domestic articles, etc., the cost is about 5 per cent of the net cost of the goods to the manufacturer. Wrought-iron grilling, office railings, and the better class of scroll and fancy work cost about two cents per pound. Wrought-iron steam and water-pipe is coated for about the same expense per pound as is required to paint it. A further benefit to this class of articles is, that the inside of the pipe

receives the same conting as the external parts. Wrought-iron I beams, channels, and other shapes entering into building constructions can be treated very cheaply; the principal expense is first cost of the furnace; the actual operating expenses are very small—fractions of a cent per pound.

The Iron Column of Delhi.*

The iron column of Delhi, India (see Frontispice), is 20 feet high above ground, 16 inches in dinmeter at the base, and 12 inches at the top, with an ornate Persian capital 31 feet in height. The base has a Persian inscription of six lines on the western side, symbolizing the deeds of the Rajah Dhawa, who reigned in the ninth century n.c. Beck, in his "History of Iron," places its erection in the early part of the fourth century A.D., but other authorities place it in the ninth century B.C., corresponding to the inscription upon it.

Early exervations to the depth of 26 feet did not reach the hottom, but subsequently it was found to rest upon forged iron beams, bedded and anchored to the stone foundations. A short distance below the ground it is 2 feet 4 inches in diameter, and evidently was forged from a large number of wronght-iron blooms. Its estimated weight is seventeen tons.

It stands alone above all other relies, a monument commemorative of the state of the mechanical arts in prehistoric times, not only for its construction and preservation, but its transportation from some unknown and evidently far-distant place of manufacture and its erection in situ. This would be considered, at the beginning of the twentieth century, an exceedingly creditable example of engineering skill, and it will probably remain centuries after most of the present-day ferric constructions have crambled to red rust.

It is free from corrosion, and while this in a measure may be due to the climate of India not being inducive of corrosion, it cannot alone be the reason of its protection, for other iron articles, both large and small, bear testimony to corrosive effects under the same exposure and climate. A reason for its non-corrosion has been given: that in the earlier days following its erection it was considered a part of the religious duty of every pilgrim to the holy shrine near which it is

left upon the column and thus protected it. But for the past two hundred years or more, so far as known, no such greased-pole gymnastical devotions have been practised, and the coatings of oil, if any ever were thus applied, must have long since been dissipated, as they would doubtless have been palm or some other vegetable oil or camel's fat, all of a non-siccative nature.

The ornate capital is as free from corrosion as the shaft of the column, so unless the pilgrims climbed this as well as the shaft (as a sailor-boy mastheads his ship's truck), and possibly stood on their heads as a further sign of exalted zeal, the capital could not have received the oil treatment to protect it. The part of the column underground surely had no such aerobatic oleaginous distribution, and is comparatively as free from corrosion as the part above ground. Every indication in the appearance of the column shows that after it had been forged and finished, the inscriptions and capital still bearing the chisel-marks on the ornaments, it was subjected for its entire length to a process identical to that of the modern Bower-Barff process, which has proven to be quite as effective to prevent corrosion in this instance as in any of the modern examples of this protective method.

CHAPTER XVII.

GALVANIZING. ELECTRO-CHEMICAL AND OTHER ANTE-CORROSAVE ZINC PROCESSES.

Galvanizing.*

Galvanizing to protect the surface of large articles, such as enter into the construction of railway viaduets, bridges, roofs, and shipwork, has not reached the point of appreciation that possibly the near future may award to it. Certain fallacies existed for a long time as to the relative merits of the dry or molten and the wet or electrolytical methods of galvanizing. The latter was found to be too costly and slow, and the results obtained were creatic and not satisfactory, and soon gave place to the dry or molten-bath processes as in practice at the present day; but the difficulty of management in connection with large baths of molten material, the deterioration of the bath, and other mechanical causes limit the process to articles of comparatively small size and weight.

The electro-deposition of zinc has been subject to many patents, and the efforts to introduce it have been lamentable failures in both a mechanical and financial sense. Most authorities recommend a current density of 18 or 20 amperes per square foot of enthode surface, and aqueous solutions of zinc sulphate, accente or chloride, ammonia chloride or tartrate, as being the most suitable for deposition.

Herman's process has been experimented with on a commercial scale, the chief feature being the addition of the sulphates of the alkalies or alkali earth to a weak solution of zine phosphate.

Electrolytes made by adding caustic potash or soda to a suitable zinc salt have been found to be unworkable in practice, on account of the formation of an insoluble zinc oxide on the surface of the anode and the resultant increase.

also constantly getting out of order, as more metal is taken out of the solution than could possibly be dissolved from the anodes by the chemicals set free, on account of this insoluble scale or furing up of the anodes, which sometimes reaches \(\frac{1}{6} \) inch in thickness.

To all intents and purposes the deposits obtained from acid solutions under favorable circumstances are fairly adhesive when great care has been exercised to thoroughly scale and clean the surface to be coated, and which is found to be the principal difficulty in the application of any electro-chemical process for copper, lead, or tin, as well as for zine, and that renders even the application of paint or other brush compounds so futile unless honestly complied with. Unfortunately these acid zine coatings are of a transitory nature, their durability being incomparable with hot galranizing, as the deposit is porous and retains some of the acid salts, which cause a wasting of the zine and consequently the rusting of the iron or steel. Castings conted with acid zinc, rust comparatively quickly, even when the porosity has been reduced by oxidation, aggravated no doubt by some of the corroding agents, sal-ammoniae, for instance, being forced into the pores of the metal. In wrought iron, the einder is porous, and holds the acid, and induces corrosion.

The relative porosity of zine coating, applied by different methods, is shown by the following micrographs, Figs. 25 and 26, taken from *The Engineer*, September 28, 1894.



Fig. 25. Zinc coating applied by hot galvanizing process, magnified five diameters.



Fig. 26. Deposit from zine sulphate solution (acid), magnified five diameters.

results except for the commoner articles of use.

The Cowper-Coles process of electro-zincing articles claims to overcome all these difficulties, and plants are in process of erection with a bath of some 14,100 gallons capacity, capable of turning out forty tons of light work per week, and in which it is proposed to treat the plates of vessels sixty feet in length upon one or both sides, and the frames of such vessels as torpedo-boat destroyers and kindred eraft after riveting up. These plates and frames are given a thin coating of zine by this process that appears to be perfectly uniform in character and extent whatever the shape of the piece may be, and however numerous the lugs, flanges, mortises, or core-holes. It is called "zinc-flashing"; that is, coating the iron or steel article, after pickling and cleaning, with a thin coat of zine about one onnce per square foot of surface, which resists the inclemency of the weather and mechanical injury as well as a thicker coat, and is found to afford sufficient protection in most cases, and is adequate protection until such time as it is ready to receive the usual paint coatings.

To obviate any tendency of the paint to peel from the zine surfaces, as it generally manifests a disposition to do, it is recommended to coat all the zinc surfaces, previous to painting them, with the following compound: One part chloride of copper, one part nitrate of copper, one part sal-ammoniae, dissolved in sixty-one parts water, and then add one part commercial hydrochloric acid. When the zine is brushed over with this mixture, it oxidizes the surface, turns black and dries in from twelve to twenty-four hours, and may then be painted over without danger of peeling. Another and more quickly applied coating consists of biehloride of platinum, one part dissolved in ten parts distilled water and applied either by a brush or sponge. It oxidizes at once, turns black, and resists the weak neids, rain, and the elements generally.

There are also a number of trade-mark, or proprietary, mixtures to prevent the peeling of paint applied to zinc. "Uniter," an English compound, and "Galvanum," an American paint in light-brown and darkgray colors, are favorably recommended. Carbon and asphaltum paint, containing a large percentage of bisulphide of carbon in the vehicle, also adheres well to galvanized iron. Its nauseating odor

Galvanized-iron sheets that are corrugated after galvanizing corrode more rapidly than uncorrugated sheets. Sharp angles and twists in the sheet also corrode quickly. The thin zine-coating atoms are brittle naturally, and are opened to allow moisture to reach the metal they cover. This being a more clastic metal, plates coated with it do not show the bending effects so strongly, yet they are apparent.

Double-coated tiu, zine, or terne plates are from two to three times more resistant to corrosion than single-coated plates. The second coating, like the second coat of paint on a painted surface, fills the shrinkage, cracks, and pores in the first coat. Galvanized-iron pipes used for gas and water service in the ground have only a life of 12 to 15 years, the outside coating of zine being destroyed by galvanic action induced by the acid elements in the soil. If the soil contains furnace einders, the corrosion is hastened. The screwed ends and other parts of the pipe where the galvanizing has been cut away are the parts first corroded. In general all galvanized pipework is so poorly cleaned from mill-scale and grease prior to galvanizing, that the pipes are less enduring than with a common coal-tar pitch dip conting.

Zine surfaces, after a brief exposure to the air, become conted with a thin film of oxide—insoluble in water, which adheres temciously, forming a protective conting to the underlying zine. So long as the zine surface remains intact, the underlying metal is protected from corrosive action, but a mechanical or other injury to the zine conting, that exposes the metal beneath to the presence of moisture, causes a very rapid corrosion to be inaugurated, the galvanic action being changed from zine positive to zine negative, and the iron as the positive element in the circuit is corroded instead of the zine.

When galvanized iron is immersed in a corrosive liquid, the zinc is attacked in preference to the iron, provided both the exposed parts of the iron and the protected parts are immersed in the liquid. The zinc has not the same protective quality when the liquid is sprinkled over the surface and remains in isolated drops. Sea air being charged with saline matter is very destructive to galvanized surfaces, form-

gaivanized-iron the-rois, angles, and other construction simples, in blind confidence of the protective power of the zine coating; else in supreme indifference as to the future consequences and catastrophes that may arise from their failure.

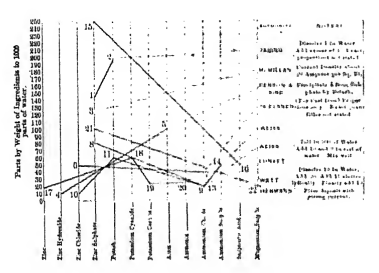


Fig. 27. Zincing solutions recommended by various authorities.

The comparative inertia of lead to the chemical action of many acids has led to the contention that it should form as good if not a better protection to iron than zine, but in practice it is found to be deficient as a protective coating against corresion. A piece of lead-coated iron or terne plate placed in water will show decided evidences of corrosion in twenty-four hours. This is to be attributed to the porous nature of the coating, whether it is applied by the hot or wet (acid) process. The lead does not houd to the plate as well as either of the other metals, zine, tin, copper, or any alloys of them. Lead-coated iron corrodes rapidly when exposed to the gases of combustion. The usual weight of lead-coated terne-plates is about \frac{1}{2} ounce to a square foot, while hot-process zine coatings weigh from \frac{1}{2} ounces minimum to 3 onness maximum, depending upon the

Description of Article,	Square Foot.	Increase of Weight.
Thin sheet iron026 inch No. 22 B. W. G	$\frac{1.76}{2.19}$ "	18.2 2.0 6.72 1.00
Tin is often added to the hot bath fo	r the purpose of	obtaining a

Tin is often added to the hot bath for the purpose of obtaining a smoother surface and larger spangles or facets, but it is found to shorten the life of the coating considerably.

A portion of a zine coating applied by the hot process was found to be very brittle, breaking when attempts were made to bend it; the average thickness of the coating was .015 of an inch.

An analysis gave the following result:

Tin	2.20
Iron	3.78
Arsenie	trace
Zinc (by difference)	94.09

A small quantity of iron is dissolved from all the articles placed in the molten-zine bath, and a dross is formed amounting in many cases to 25 per cent of the whole amount of zine used. This zine-iron alloy is very brittle and contains by analysis 6 per cent of iron, and is used to cast small art ornaments from.

Nickel coatings produced galvanically will not protect iron from corrosion unless .02 inch thick.

A hot galvanizing plant having a bath capacity of 10 feet by 4 feet by 4 feet 6 inches outside dimensions, and about 1 inch in thickness, will cost \$625, and will hold twenty-eight long tons of zinc, which at four cents per pound will require \$2500 to fill it; the heating of this mass of metal and its ever-changing cold immersions, with the waste by dross and extra thickness in spots, is a constant source of annoyance and expense.

The cost of an electro-chemical or wet-bath Cowper-Coles plant of 6700 gallons bath, size 30 feet by 6 feet by 7 feet, will be but slightly more than the hot bath given. There is no dross formed galvanized from wire, as per lottowing faine:

RESULT OF PROCESS TEST MADE ON SAMPLES OF CHARGON THOS WHITE CONTROL WITH ZING BY VALUE & PROPERTY.

Process Used to Cont the Ware.	Grains of Zin Por Sopiaso Tout,	Objice Per Signar Lie F	Number of times uniouse times, manager beauti and out blowing that allow topper
Hot galvanized	116 1	1 18 1 02 1 26	1 3
September and the contract to the second description of the contract to the co			í

A Cowper-Coles process bath of a capacity of about 4000 gallons will treat ship-plates 18 feet long, and will require an electrical energy of 2000 amperes of 5-volt electro-motive force.

With equal amounts of zine per unit of area, the zine conting put on by the cold process is more resistant to the corroding action of a saturated solution of copper sulphate than is the case with steel conted by the ordinary hot galvanizing process; or, to put it in another form, articles conted by the cold process should have an equally long life under the same conditions of exposure that hot galvanized articles are exposed to, and with less zine than would be necessary in the ordinary hot process.

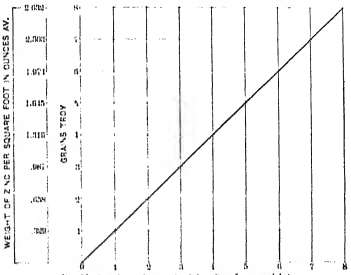
The hardness of a zine surface is a matter of some importance. With this object in view, aluminium has been added from a separate crucible to the molten zine at the moment of dipping the article to be zinced, so as to form a compound surface of zince aluminum, and to reduce the waste formed from the protective coverings of sal-annuoniae, fat, glycerine, etc. The addition of the aluminum also reduces the thickness of the conting applied.

Cold and hot galvanized plates appear to stand abrasion equally well. The thickness of the conting being the same, tests by means of the Schlerometer show; cold galvanized sheet, 6; hot galvanized sheet, 6; terne-plate, 2; tin-plate, 2. The figures represent the load in grammes upon a diamond point, just sufficient to cause it to scratch the specimen.

The attempts to electro-zine iron and steel wire for wire standing

Both pickling and hot galvanizing reduce the strength, distort and render brittle iron and steel wires of small sections. Zinc fuses at 775° F, and vaporizes at 830° F. Hence the necessity of the salammoniae bath that covers the molten zinc, prevents volatilization and acts as a flux to unite the zinc and iron. The bath is usually kept at about 1000° F. Steel wire of high breaking strain has its

TABLE GIVING THICKNESS OF ZING REQUIRED TO WITHSTAND VARYING NUMBER OF IMMERSIONS IN A SOLUTION OF COPPER SULPHATE.



No. of I minute importains in extended equiton of copper sulphate

Fig. 28,

hardness, and consequently its ultimate tensile strength and clongational efficiency, reduced by drawing the temper and the formation of an iron zine alloy on the surface of the wire by as much as from 5 to 10 per cent. It is the practice when coating steel wire to keep the bath at as low heat as possible and to run the wire through it at a bigh rate of speed. Both these coverations lead to a waste of

The Cowper-Coles or cold-galvanizing process is used for the purpose of zincing the skin plates and frames of the torpedo-boats and torpedo-boat destroyers built for the English mays. A plan and elevation of this plant is given in *The Engineer*, Feb. 28, 1894.

of rendering it worthless.

The industrial importance of the successful application of this cold-galvanizing process can hardly be overestimated, even if its application is only to the nurine constructions of the future, and it is found to be in any degree inapplicable to our present structures and vessels in use. The permanency, continuity, strength, and density of the conting given by this process is in all respects count to that of hot galvanizing, and the thickness of it can be made superior to that given by the hot. Considering the success that has attended the use of zine to prevent corrosion in marine boilers, where concentrated hot saline fluids are the excitant medium, nided by the electrical conditions attendant upon the combustion of large quantities of fuel, it may not be considered a wild prophecy to expect that with all of the internal metallic parts of a steam vessel protected by an application of zine plates secured to the framework of the structure similar to the application of zine to marine boilers, that these plates may receive the energy of corrosion, and if not neutralizing it entirely, at least pass it along in the form of a deposit to convenient pockets, where it could be removed, the same as is now done with the washings and dirt from the fire-room bunkers and ballast-chambers. This internal electro-chemical process of protection does not

ings and dirt from the fire-room bunkers and ballast-chambers.

This internal electro-chemical process of protection does not appear so chimerical as at first one might suppose. Dr. Henry Wurtz* has proposed the protection of mining plants subject to the intensified corrosion due to the decomposition of pyrites and other minerals in the mine waters, by connecting all of the metal partions of the mine as the negative elements with a dynamo of sufficient force to overcome the strength of galvanic energy due to the surfaces exposed being excited by the corrosive liquids in the mine, the positive terminal to be connected to a mass of hard coke in the mine sump. These conditions vary but slightly from those existing in the ship, and it is not improbable that experiment will determine that both these systems could be made to work successfully.

an bodies, and set up volude action in all cavities, fissures, seams, and contact surfaces in the metal, which, though slight and not easily detected, will in time calarge and waste them away sufficiently to sap the strength of the mass.

Metallic salts and acids in mine waters intensify the corrosion of all metals exposed to their action. The metal work of railway tunnels is also disastrously affected by the condensed vapors of sulphur, carbonic acid, and the ever-present moisture due to such locations. The corrosion of the metals decreases the resistance of the water to voltaic circuits, this corrosion by liquids being voltaic phenomena in all cases. In many cases it is intensified by the moisture being in the form of drops instead of being uniformly spread over the whole surface.

Acids and acid salts which are capable of taking up iron oxides into solution still further enhance the destruction by removing such oxides and exposing the surfaces of the metal to a fresh attack of the corrosive element. The saline matter in solution that excites voltaic action need not be acid. Any neutral salt which decreases the resistance of the water will qualify it to act as the necessary liquid medium of a voltaic circuit. Sea-salt is the commonest of all such neutral salts, together with the other chlorides and sulphates of sea-water. It enables corroding voltaic action to be set up on all ferric bodies immersed therein or in the air impregnated with their substance.

neutral salts, together with the other chlorides and sulphates of seawater. It embles corroding voltaic action to be set up on all ferric bodies immersed therein or in the air impregnated with their substance. The Journal of the Society of Chemical Industry, London, February 28, 1894, details some experiments with the galvanic action of seawater upon iron and steel structures in various relations with each other, such as the constructive parts of trusses, boilers, etc., to prevent the corrosion for which the use of zine and other easily oxidized metals and alloys are suggested, and to be so placed and connected to the structure that they will form the electro-positive element of the ever-present galvanic circuit, and by their decomposition protect the structure. Mr. D. Phillips, in a paper read before the Institute of Civil Engi-

neers, in 1885, cited the result of an experiment, where "surfaces of bright pieces of plate iron, immersed in cold sea-water for over ten years have been thoroughly protected from corrosion by the aid of pieces of metallic zing in metallic contact with the iron; while a simi-

whatever. The water was changed twice annually, and the oxide removed from the zine by filing. Under these circumstances the iron became gradually coated with a film of leader-colored deposit when wet, but hard and white when dry. The effect in other respects was that, on every occasion that the oxide was removed from the zine and the deposit from the iron specimens, on being returned to the water small globules formed on the zine, and on reaching $\frac{1}{100}$ inch in diameter released themselves and flow to the surface."

The proportions necessary to insure complete protection from corrosion in marine hollers are one square foot of zine to fifty square feet of heating surface in new boilers, which may be diminished after a time to one in seventy-five or even one in one hundred square feet. Merely placing the zine in trays, hangers, or strips will not insure metallic contact. The better and generally recognized method of fixing the zine is to place a number of study in the sides of the furnaces and combustion-clambers, and to bolt on to these study the zine plates, which should be about 10" × 6" + 1". It is important to see that the contact surfaces are clean and bright, and the nut screwed close down to the zine to exclude the water and deposits from the contact surfaces, thus comparatively insulating them and preventing the galvanic action. Otherwise the zine is acted upon mostly as a solvent that renders the water innocuous or non-exciting, but does not prevent the water from forming a hard scale when it is saturated.

Sheet zine has proven to be a durable rooting material. Zine is reduced in density from 6.86 to 7.2 in the process of redling into sheets, which closes the pores and renders the metal less affected than tin-plate from the ammonia, carbonic acid and atmospheric gases.

Berlin zine roofing-plates (unpainted) have been found to be not materially affected after many years' exposure. The weather formed a thin film of oxide on their surfaces that effectually prevented further oxidation. A few cases reported are as follows.

The Cloisters of Canterbury were covered with zine roofing and were uninjured after 33 years,

The Portsmouth Dock Yard Buildings' roofs were uninjured after

of flaxing or non-drying oil that fills the pores of the tin, and if the roofing is painted soon as laid, this film prevents the paint from adhering to the tin, just as a machine grease prevents a paint from bonding to a surface. A few months' exposure to the atmosphere slightly oxidizes the tin, and this oxide absorbs the oily coating and allows the weather to wash it away and the paint has a clean metallic surface to bond to. Tin-plates doubly dipped are less porons and more durable.

The quality of commercial tin-plate is greatly inferior to that made forty years ago, and appears to retrograde yearly. Lead, antimony, and other metals are mixed with the tin in the dipping-bath, and greatly reduce its resistance to corrosion. None of the adulterants form a true alloy; they are only mechanical mixtures. They all differ in oxidizing power and electrical affinities. The lead is electronegative to the tin and zinc, which again are of opposite electrical natures.

The amount of sulphurous and carbonic acids and ammonia in the atmosphere is enough to form the excitant element needed to decompose them one after another, until the coating is made porous and the iron is corroded in turn. The life of the tin-plates is also governed in a great measure by the want of care that they should receive in the preliminary pickling with muriatic acid, to free them from the mill-scale that always attends their rolling. Ordinary washing with line-water does not remove the whole of this acid, and the tin conting usually has a double galvanic pile in a sandwich form, ready for duty on the least encouragement.

There are brands of tiu-plate as honestly coated at the present day, and as reliable in all respects, as any ever made, but they are an exception, not the rule. Price and the gallibility of the purchaser govern, as in many other modern industries. (See Fig. 6, page 38.)

Red-lead paint contings soften tin roofing, but do not wholly destroy it, although some of the tin may be changed to a white oxide that is easily removed by atmospheric influences.

CHAPTER XVIII.

INERT PIGMENTS, OR ADULTERANTS.

THE different substances known as inert pigments are used to a great extent in the preparation of nearly all mixed paints, particularly in the house paints, where the amount of one or more of them frequently exceeds that of the base pigment.

However admissible their use (on account of cost only) may be in paints not classed as protective ferric coatings, their durability in any case is determined by the character of the weakest element in the associated group to resist atmospheric conditions, whatever the base pigment may be.

The manufacture of "patent pa uts" would be almost nil were it not for the very liberal use of these inert pigments. They are said to correct almost every detrimental quality in the basic pigments. Yet with all of their boasted virtues, there is hardly a manufacturer of paint willing to admit their use, or that will furnish an analysis of his product that contains them.

Many of the uses and characteristics of these ments, fortifiers, or adulterants have been mentioned in the basic pigments chapters and elsewhere in this work, but are brought together here for comparison and ready reference.

Carbonate of lime (CaCo) in some form other than as quickline (calcined limestone or marble) is often used as a desirable adulterant of many paints. It is claimed to be specially favorable to correct the sulphur element present in iron exides.

Chalk is a friable carbonate of line that, on account of its cheapness and several colors, is the most used. Its specific gravity is 2.2 to 2.8. According to the basic oxides in it, the relors are white, red. reaction between the oil and itself that results in the formation of a lime soap, which is not at all a durable substance.

Putty, however (a mixture of whiting and oil), is a very durable body, and withstands atmospheric exposure and water remarkably well. In this form it illustrates the theory that the pigment is the life of the paint. The small amount of oil in the composition of putty is the cause of its quick drying. Its mass, when applied, greatly exceeds that of a paint coating, and its shrinkage solidifies, instead of rupturing it, by a movement in a number of directions, as in a paint.

Baryles (Heavy Spar), Ba.SO₄. Specific gravity, 4.3 to 4.7. The natural sulphate of barium, consisting of one atom of barium oxide (BaO) ~ 65.67 per cent, and one atom of sulphuric acid, 34.33 per cent. It is the heaviest of all minerals, and is found in all stages of purity, in transparent, colorless, white to yellow crystals, also in a granular and compact form in heavy beds resembling marble. It is common in all metallic veins, allied with, or changed to, calc-spar, spathic iron ore, cerussite, quartz, limonite, pyrites, and other substances. It is the white variety that is ground for a pigment, but lacks the opacity or light-reflecting or coloring power to form of itself a good pigment. It grinds hard, splintery, and irregular, and is used to give weight to paper-stock, zinc oxide, gypsum, and all the other light pigments that lack weight to enable them to masquerade as white lead. (See Chapter VI.)

Barytes brightens light-colored paints, though of poor coloring or light-dispersing power; also spreads easily and saves oil. It is mixed with nearly all pigments, and by the use of a stiff or short bristle brush, covers a large surface with a resemblance of a good paint.

White lead does not cover so well with barytes, but zine oxide covers better. Zine oxide lacks weight that barytes furnishes and also saves oil, advantages not ignored by the cheap paint-compounders. Barytes does not unite with the oil in any degree. From

bonate with sulphuric acid and precipitating the artificial barytes from the solution. It is less exystalline than the natural subpliate and has a greater covering power. Baryta white, permanent white, constant whate, etc., are of this class of pigments. Blane Fixe mixed with the mineral barytes compose the principal substances in most of the commercial white

off in water to give a finer product.

of volutiles or quick driers. But yies along is the product of jugments. Floated barytes is the ground natural sulphate of baryta floated

Artificial barytes (Blanc Fixes) is made by heating larinin car-

putent paints. Lithopone, a trade-mark for one of these mixtures, has an extended sale under the guise of white lead. Trade marks are easily invented,

but they add no durability to the paint. They move around in paint literature as easily as some of the substances covered by the name

move in the vehicle that gives them a home, it not rest Barytes as a pigment, exposed to air or on underground badies, condenses water and earbonic acid and is converted into a carbonate with the evolution of sulphuretted hydrogen. This decomposing feature in barytes seems to be ignored by paint compounders, but to it the failure of many coatings can be attributed.

Rose's experiments show that burytes in any form, when acted upon by water, evolves sulphinetted hydrogen and sulphinions acid, leaving the decomposed lime free,

Hansfeld has also shown that sulphate of lime is decomposed by the galvanic action of two metals or metallic oxides in contact under the ordinary exposures of a paint. When both barytes and gypsum are present in a paint, this galvanic action between three substances

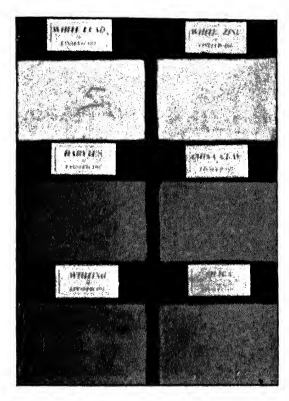
is certain to occur. Mixtures of burytes and gypsum with the oxides or carbonates of zine or lead will in no degree protect any one of them. They decompose one after the other; the first to break down only adds to the electrolytic energy to harry up the decomposition

of the others. The thin film of oil in which the pigments are embedded is sufficiently porous to admit the atmospheric moisture and carbonic acid necessary to start up the disintegrating process. Putty made from barytes, whiting, and oil dries into a hard,

limitale summer along the

temptation to use it in place of these pigments is not always resisted. As a rule no responsible paint firm with a business reputation to sustain will sell a barytes adulterated paint under their name.

Additional data about the presence of barytes in a paint is given in Chapter XXIX. The covering and coloring power of barytes in comparison with white lead and zine oxide are shown by Fig. 29.



1 p. 29 Covering power of mert pigments.

Brick-dust is used to a large extent to adulterate red lead and other red paints; even the low-price iron oxides do not escape it, particularly with other proportions of the same ingredients, whether applied to wood or iron. In general the tiles added nothing to the quality of the paint, only reduced the cost of it. They are practically movidizable by atmospheric influences or weak acidulous solutions, and are electronegative to metals or their oxides. In any electrolytic action set up by any cause in a paint conting of which brick-dust is a part, the tendency will be to decompose the other pigments, possibly, before electrolysis is developed in the covered iron surface. (See page 53.)

Feldspar. Specific gravity, 2.5 to 2.8. Decomposed nica, gradite, gneiss, and most forms of basalt form this class of adulterants, and are all inclined to further decomposition on exposure to the weather. Many of the fire-clays used in the manufacture of fire-brick are broken down and decomposed feldspars. Its use in the composition of a pigment is of the most unreliable character in all respects. It is as poor a substance for an adulterant as nature furnishes. Mixed paints frequently contain 15 to 20 per cent of it. Feldspar carries a large amount of water loosely held and frequently acidalated, also sand, etc. It is easily whipped up in the oil and mixes well with graphite for dark-colored paints.

Gypsum (Sulphate of Lime). Natural mineral (hydrated), Catt.SO₃ +H₂O; calcined, CaO.SO₃. Specific gravity, 2.4 to 2.8. The untural mineral ground is the plaster that farmers use on their crops to attract and condense atmospheric moisture. Calcined to expel the one atom of water held in its natural state, it becomes the common plaster of Paris, used for the hard finish of plastered walls of buildings. The process of grinding is supposed to drive off the one atom of water it holds naturally, by the heat developed in the dry grinding-will, but this is soon replaced upon a short exposure to the atmosphere, and when used for the hard finish, it must be heated again to dispel it, or a porous wall conting results. A hydrated sulphate of lime contains over 18 per cent of water.

When used in the composition of a paint, it must be thoroughly calcined, and is so specified by parties who allow its use. Neglect of this carries the moisture into the paint, where some portion of the sulphur element in the gypsum is released, and combining with the

right cause, viz.: too much of a sulphurons adulterant.

Gypsum grinds easily, is opaque, and incorporates readily with

Gypsini grinds easily, is opaque, and incorporates readily with most pigments and the vehicle. It is not liable to set or settle in the paint-bucket or package, and probably is the best of all the inert substances to use as an adulterant.

The extra atom of sulphure in the natural mineral other than that meessary to form the sulphurous-acid compound is strongly combined, but if the mineral or ground pigment is calcined at a temperature higher than that necessary to release the one atom of water, the sulphuric-acid atoms are excited to a degree that will afterward manifest the same destructive properties as the same element does in any other pigment or substance, and as noted above in the "livering" of the green-paint coating.

A synthetical subpliate of lime is supposed to be formed when iron ore is rousted in a furnace in contact with a quantity of carbonate of lime. The roasting process, besides driving off the moisture in both the iron ore and carbonate of lime, and a part of the sulphur in the iron ore, excites the remaining atoms of sulphur to leave the ore, and combines with the now anhydrous carbonate of lime, CaO, and forms the anhydrous sulphate of lime, CaO, SO₂, described above. The process is an unsatisfactory one, as the carbonate is generally added in great excess of the amount needed to effect the chemical combination with the sulphur to form the sulphate. When the roasted iron ore is removed from the furnace to be ground, the sulphate is not distinguishable or separable from the uncombined carbonate of lime, and both are ground with the ore and appear as adulterants, that may be 5, 10, or 20 per cent, or as much as can be unloaded upon the consumer.

In whatever amount these lime products are present in the ironoxide pigment, they are both, like the oxide, anhydrous, hygroscopic, and readily attract moisture, frequently 5 per cent in oxide pigments that have been made for some time.

The synthetical sulphate of line formed in the roasting of copperas, as described in the preparation of that substance for an ironoxide pigment, is of the same character as that from roasted iron any permanency of color of protective quantities.

Kaolin (American Terra Alba). A clay of the same class as pipeclay, China clay, potters' clay, etc. The reddish color of the latter
being due to the iron and other metallic exides. Specific gravities,
2.58 to 2.76.

Their general composition is as follows:

Substances,	Paper dax White	Patter and the Light and United	Terra Allia, White and thay,
Alumina	72 23 " 65 49	Permanger 2d 25 to 21 28 7d 3d 2 61 95	Theorem ages 25 Ad to 15 Ad to 34 22 Pt 65 22 Bt 22 1 Ad
MagnesiaOxide of ironAlkaline cartls	****	1 26 to 2 51 7 26 % 1 75	1 17 " traces 1 70 " 2 05
Sulphate of lime Moisture	12 (8) to 1 (8)	1 72 7 traces	8 58 66 (0.42)

All of the pigment-clays grind greasy, and are as easily broken down and decomposed by the weather as any of the clays in building brick or mud from a mill-pond.

Mixed with tule, the clays are supposed to add some advantage to pigments of a granular character. What that advantage is the author has never been able to ascertain, but he knows they cause paint to peel or crack.

Marl. Specific gravity, 2.4. It is composed of:

Q11:		****	\$ 88 E.	cent
Silica				**
Ahunim		32	**	* *
Oxides of iron and manganese,		. 100	••	**
Water		. 4	**	14
		100		

Its gray color prevents its use as an adulterant of the white paints, but in the tinted colors it is used quite as freely as kaolin or chalk. It is difficult to pulverize on account of its greasy nature. It saves

Ochrc. A yellow clay containing from 8 to 15 per cent of water loosely held with large amounts of sand, also marked quantities of iron oxide and sulphur. When moderately heated, the lower grades of ochre contain sulphur enough to change their color to a pink or low red.

The common grades were formerly used as a coating for tin roofs. They were always subject to blistering, from the large quantity of water they carried into the paint. It is an adulterant without a single inert element in it, and its presence in a paint is generally accompanied by as poor a quality of oil as it is a pigment.

Silica (Si.SO₂). Specific gravity, 1.9, 2.5, 2.8. Is a sulphate of silicon containing one atom of silicon combined with one atom of sulphurous acid. It is found in crystals of different degrees of translucency, and forms a component part of all metallic ores; of iron ore, frequently, 50 per cent. (See Analyses of Iron Ores, Chapter 111.)

In its natural form it is one of the most imperishable of all minerals. It grinds hard and splintery, and is difficult to reduce to the fineness required for a pigment. Manufacturers of silica products subject the crystals to a bright-red heat and quench them in water, causing them to fracture and grind more easily. However, the calcination drives off a part of the sulphuric acid and renders the silica caustic, the latter condition is not a favorable one for any pigment, inert or basic.

Silien is not affected by sulphurous gases, acids, or alkalies. Floated silien or silex makes an excellent wood-filler paint. All siliens are difficult to hold up in oil, and on settling, cake very hard.

Sand, generally supposed to be the same substance as silien, is, however, of quartz formation as an oxide of silicon, specific gravity 1.44 to 1.76, only about two-thirds the weight of silica. It grinds hard and splintery, or in an irregular crystalline form, and is difficult to grind to a pigment.

Neither silies nor sand mix with other pigments, except in a purely mechanical manner, differing according to the specific gravities of 92 INERT PIGMENTS, AMOUNTS MANUFACTURED, ETC,

turer of paints will acknowledge. The covering power of silien is shown by Fig. 20.

Fine sand is used for an application to green paint on exterior surfaces with a view to affording an extra protection from atmospheric effects. It in all cases hastens the decay of the paint. It holds the moisture, dust, and other organic substances, and their easy and early decomposition results. Physics form more readily under a sand-coated paint than with a paint alone.

Tale (Steatile or Soupstone). Specific gravity, 2.65 to 2.8, Grinds greasy and flaky, is inclined to cause a paint to peel, and is repellent to the oil. Its use with knohin has been given. It is used for a special adulterant of flake and other graphitic carbons (see Chapter XIII). As a pigment it has no qualities whatever. As an adulterant its function is to enable some objectionable substance to attempt a mission that could be better performed by a straight pigment.

The above list does not exhaust the substances known as adulterants or miscalled "inert" pigments. As protective coverings for ferric bodies, the protective effects of the inert pigments in use with the basic pigments will be noted in the paint tests made to determine their value (see Chapters NXIN and XXN). Their covering or coloring powers are shown by Fig. 29.

The following amounts of inert pigments were produced and used in the United States (averaged for the years 1898 to 1901):

Barytes, all grades, 124,000 short tons. Cost of the crude numeral, \$3.30 to \$3.50 per ton.

Imported barytes, 1400 tons, including some floated. Cost of the manufactured article, \$10.50 to \$11.00 per ton.

Feldspar mined in the United States for all purposes in 1898 to 1901 averaged 27,280 tons and cost from \$3 to \$6 per ton

Ground slate and shale for pigments averaged 4.700 short tons, value from \$9.50 to \$10.00 per ton,

Of soapstone ground for pigments and foundry use, there were produced 9000 tons yearly. Value S\(\frac{1}{2}\) to 0 cents per pound.

CHAPTER XIX.

SPIRITS OF TURPENTINE.

The composition of spirits or oil of turpentine is $C_{10}H_{10}$. Specific gravity, 0.86 to 0.88, with a boiling-point always near 160° F. The several varieties of commercial turpentine obtained from the sap of firs and pine-trees are more or less viscid solutions of resins in a volatile oil, the proportions of these constituents varying according to the source and age of the turpentine-tree. Some kinds are clear and homogeneous; others are more or less turbid, holding in suspension granulo-crystalline masses, which gradually settle to the bottom, and are known to painters as "drops."

Spirits of turpentine is the product of the first distillation of the crude gum, and consists of about one-third spirits and two-thirds water. It requires about twenty-five barrels of crude gum to make two barrels of the spirits of turpentine, that when redistilled is known as refined or oil of turpentine.

The principal supply of turpentine is obtained from the American long-leaf yellow pine-tree, *Pinus palustris* (*P. australis*); also from the loblolly-pine, *P. tada*; all products of the southern part of the United States, where the Conifere are the principal trees. There are many varieties of the Conifere, and all yield gams available for distillation into turpentines and resins.

Turpentine consists chiefly of a hydrocarbon oil (C₁₀ H₁₀) and a resin called "Colophony" (C₂₀H₁₀O₂). Specific gravity, 1.07 to 1.08. It softens at 155" to 175" F, and melts between 194° and 212° F. The spirits of turpentine constitutes about 17 per cent of the yellow pine-tree sap or crude gum. The Maritime pine furnishes about 24 per cent of spirits of turpentine.

The exided gum from all of the turpentine-trees is a yellowish, opaque, tough mass, brittle and crumbly when cold, crystalline in the interior, and of a characteristic taste and odor, a distinguishing

P. nigra, and P. rotandata.

The English, from the American or Carolina Priors australis or P. twda.

The French, or Bordemy, from the Prins maintima, resembles the American turpentine in appearance, odor, and taste, and is considered to be the quickest drier.

sidered to be the quickest drier.

The Strasburg is the product from the Almes pectimate and from the spruce fir, Alics excelsa.

The Venice is the product from the Terebenthine venity, or the larch, Larix curopea.

The Hungarian is from Pinus pumilio.

The Carpathian, from the Pinns vember, has a latter taste.

The Cyprian, Syrian, or Chio, obtained in Chio, as to in the Pishwin terebinthus.

Templin, or pine-cone oil, is furnished from the corner of the Pinus pumilio and the Abics pertinuta.

The Canadian oil or Canada babain, from the Abics bulsaimed (Balin)

of Gilead), furnishes the whitest and purest of all of the impentines.

Related to the true turpentine-orle are the two volatile orle of the

conferous plants—oil of juniper from the Juniperus communis, and the oil of savin from the Juniperus salana.

A characteristic feature in American turpentures is that they polarized to the right, while most of the turpentures from other sources polarize to the left.

The crude resin from which the oil of turpentine is distilled has a specific gravity of 0.95 to 0.98, according to the time of its collection, whether in the first, second, third, or fourth year after the tree is boxed, or during the time of collecting the dried sap in the first flow in the spring, or the summer, or later in the fall. Also its freedom from sand, leaves, bark, and dirt; all of which are readily absorbed by the sticky, drying sap, and are only removed in the process of distilling the gum for spirits of turpentine. Another distillation is required to produce the oil of turpentine or "turpe" of the painter.

Fig. 30 shows the old method of boxing the trees to collect the

crude resin.

In addition to the exuded gum from living trees the collect the

ling the cord-wood, are cut down and distilled in a kiln or oven similar to that used for the production of charcoal from hard wood.



Fig. 30 Boxing the turpentine-tree.

A cord of fat pine wood yields by kiln distillation, according to the amount of the pitchy matter in it, whether it is body wood or from the limbs, tops, or decayed wood, the following products: acids. The effect of the use of turpentine in a point or varnish is to flatten the gloss or lustre.

Even with a pure turpentine not more than 31 per cent is admissible in a paint; and less than this if the turpentine is poor or fatty.

Pure turpentine-oil is adulterated with condc or undostilled turpentine, light-colored resin-oil, and resin.

These adulterations are detected by the pyredigneous smell and nauseous after-taste on the tongue and by the change in the specific gravity. Also by the reaction produced by adding 8 drops of strong ammonia to 90 c.c. (1.422 cubic inche to the turpentine. The following are the results:

Marie - 1, 14 - 26 P		
Pure Oil of Turpentine.	Specific litarity	Then trems mitte Ammieria.
Pure oil of turpentine recently distilled	0.8678	Nuclied, The intpentine evapointes quickly, No residuina,
Old pure oil of turpentine 7.4534 pounds per gal.	0 8693	Solubtics in a few seconds, forming a white crystalline substance with the consistency of butter.
Pure turpentine with 10 per cent of resin spirit	0.8784	torms an emulsion, who is rapully be- comes clear. The moments which separates has a pale vellow color.
Pure turpentine with 10 per cent of unlistilled turpen- tine. 7.3496 pounds per gal.		forms an emulsion which becomes clear on standing, gives a semi trans- parent sediment of a blaish color, the liquid above being colorless.
Pure turpentine with 10 per cent of resin	0,8831	Each drop of annuous appears to solulify as it falls into the oil. On agitation the whole solidates into a consistent transparent transp.

Characteristics of Oil of Turpentine,

Pure oil of turpentine has the composition of $C_{\rm in} H_{\rm 160}$ and at a specific gravity of 0.839 weighs 7 pounds per gallon. At 60°F, the gravity is 31° Baumé, and it weighs 7 pounds per gallon. At 60°F, pure turpentine should weigh not less than 6.802 nor more than 7.278 rounds per gallon.

blotting-paper filter.

An average quality of turpentine boils at 320° to 350° F. and has a flash-point of 103" or 104" F.

Crade turpentine resin boils at 316° F.

Crude turpentine resin, specific gravity 0.98 to 0.95, is dissoluble in water, but readily soluble in other or spirits of turpentine and in six parts of alcohol. The alcoholic solution has an acid reaction.

Bromine and iodine act violently upon it. When brought into contact with a mixture of nitric and sulphuric acids it takes fire. Turpentine is a solvent of all oils and resinous gums at ordinary temperatures, but some of the fossil resins require a low heat to aid its action.

Adulterants of Turpentine.

Kiln-distilled spirits of turpentine contains pyroligneous and other acids, specific gravity, 0.80 to 0.84.

Crade petroleum, specific gravity, 38° to 48° Baumé; weight, 7.00 to 6.62 pounds per gallon.

Benzine, specific gravity, 54° to 62° B.; weight, 6.39 to 6.10 pounds per gallon.

Naphtha, specific gravity, 62° to 70° B.; weight, 6.09 to 5.79 pounds per gallon.

The pyroligneous acid in turpentine distilled from dead-fat pinewood settles out partially after standing, but commercial brands not redistilled still contain some amount of the acid.

Commercial spirits of turpentine has a specific gravity of 32° Baumé. Any addition of petroleum of 40° B, will be shown by the rise in the hydrometer; each 3 to 5 per cent of petroleum added causes a rise of 1° B, on the scale. If the adulterant is benzine or naphtha, then the difference in the specific gravity is very marked and will at once determine the character of the adulterant, it being much lighter than coal-oil or kerosene.

For the detection of resin in the spirits of turpentine, the polariscope-test is the only one that can be considered strictly accurate, but it is a delicate one, and requires experience to determine results.

The structure of the st

a test-stabe and notice the precipition of the most of the resin normal in all turperates the given of the few will be found on top of the third in the table. The second of the distributions consists in heating the second of t

acid one part and four part part or three to the in

The Journal of Chemical Lie and No. 1 No.

The U.S. Navy Department to a second second of the Association of a piece of white expectation is a second with a finite expectation.

light, if the turpentine is pure and sold as the control of the control of

after 5 to 7 minute. A fauticie element in the control of the cont

miles of forest to supply the temperative product \$8,000,000 to \$8,000,000 yearly. Early is a re-increasing from 5 to 8 per cert. The results of temping is decreasing in more than the result of the results of the results amounts.

The unit of product for a respective as a function of 2500 trees from 100 to 300 messages as a second size of the trees, or an expense of 1 messages as a second such as a second one half a cord of wood product as a second second size tree as a second sec

amount, if it all could be redirected and the second of th

redistill to about 10 vallon of the ed as a grant of the

about 25,000,000 callon of turpentine, or a little more than the present (1905) requirement, if the supply came from this source alone.

When the lone lest pine force thave practically disappeared, they will have to be criefally elemed once and for all in order to produce a quantity of turpenture equal to the present demand for one year.

A barrel ('40 to 260 points) of the crude turpentine resin, when distilled, yield strong 10 to 11 gallons of turpentine spirits that need to be redictibed to afford a pare oil of turpentine. About one half or five eighths of a barrel of 6 in (170 to 190 pounds) is also a result of the sin tilliation. There is in it remailed for regin oils of a number of state, since a perite gravity, range from 0.960 to 0.9910. It also turns here to ear receive educate of commercial resins; those known as W. W. excited state, W. G. evandow das a are the finest and most variable, being produced from the interest run or virgin cap. The first are expected as a contraction of five year. That the trees run rean, an interest quality, produced, that is graded X, (very clear); then W. L. K., J. H. to A., the factor being after to black, and rated continuously as a rate, produce that the

The flow of the angle of the VI steved or virgin cut free is from that to Bottlemed on 200 and 600 points, for the first very, and requires 100 to 200 points of the flow of the flow of the flow of 60 barrels in the fearth of a 20 state. The flow of the pooler chale of crude resin, that contains that the forther bart to the pooler chale of crude resin, that

The solver of the effective endiner to point or xamilla is to form the percendence in the solver to the artifact renders them non-drying association of the solver.

Therefore be a close of the restriction of it took in the bounds, indeed to the restriction of C_i , H_i , O_i with see, and enumed be properly a constant count to the form the restriction of the restriction of the restriction of the first constant to the life of the point count C_i of C_i .

Adulterations of turpentine with resin oil residue left after evaporating a small quantit sticky nature and resinous odor after it is ignite. The use of tank cars that have been used to

leum is responsible for a great deal of the ung The crude oil of turpentine thus transported as will carry over enough of the petroleum to see gravity of the turpentine.

From a large number of tests of commercial State associations of painters, and by individual menters, the general result appears to be the samples showed adulterations ranging from 5 are no penalties for the adulteration of either oil, and when adulterations are present in any erally, the detection of them is beyond the

The Secretary of Agriculture for the Unite 1890 reports that at the present rate of consetthe long-leaf or turpentine pane will be exhausers. Practically the yellow-pane forests of Neuro exhausted, and the production of turpen centred in Georgia, Alabama, and Horda

purchasing agent or minter.

SPIRITS OF TURPENTIL

2,098,810

65,000 gallon

22,000,000

1,600,000 barre

States for the year 1897, to only six of the E as follows:

THE PROPERTY OF THE PROPERTY O		
Germany.	2,418,790	44
Italy	398,710	**
Netherlands	2,359,590	**
Great Britain,	8,476,700	44
Total.,	15,817,600	**
Other countries,	682,100	44
	16,500,000	**
United States consumption	5.500,000	**

Total production.



rich in pitch. They contain as hi 21 per cent is turpentine. The latt

point (150° F.) as that from the nort other properties. The kiln products

charcoal, pitch, etc.) from one butt had a value of \$275." The tree g where the spruce, hemlock, or any Co after utilized for its turpentine-appel Columbia to Mexico in large forests, t

in height. The bark is useful for tand

CHAPTER XX.

CARBON BISULPHIDE

(CARBON DISCLEHIDE SULPHO-CAR This carbonic anhydride (CS₃) has a spec

1.027 to 1.072. At 60° F., 1.272 or 10.6135 pecific gravity of its rapor at 60° F. is boiling-point of the continercial article is 118.

109.4° F.

It is composed of 15.8 per cent of carbon an (CS_2) , and is produced by passing the vapor phurous-neid gas, $S(\ell_2)$ over charcoal kept at commercial method of manufacture. It is

vapor mixed with air takes fire at about 30 great violence. It is a colorless, heavy, very van acid, pungent taste and a very fetid allie

oils.

It is used to extract the oil from seeds, particularly linseed, which is heated and pressed to remove some of the oil before being submitted to the action of the bisulphide. The residue cake contains only 2 per cent of oil and about 7 per cent of water, while the cake from the ordinary process of manufacturing linearly of from the steamed linseed contains 9 per cent of oil and nearly 15 per cent of water. The oil so expressed is of good color, but contains more mucilage and less of the glyceride clement.

The loss in the manufacture of the bisulphide is about 50 per cent of the charcoal and 17 to 18 per cent of the sulphin.

The use of the bisulphide of earbon for a paint vehicle is more for the cheaper grades of roofing or color paints and for whoden structures of minor importance than for the better grade of house or ferrie paints, though in many of the latter it is used freely, judging from the odor. Its use is simply as an adulterant of the oil and to cause a quick drying of the paint, and wherever used it may be considered to accompany a cheap oil, and the grade of the pigments mixed with it will in general be as low as the vehicle.

Like benzine driers, it sensibly lowers the temperature of the surface of the body being covered, and in cool or damp locations this reduction is often enough to reach the dew-point and cause a sweat-deposit on the surface of the paint, causing it to peck

There are special grades of earbou blacks, or brailphide of earbou paints, under many trade-marks, specially noted in trade literature for their excellence as coatings for brine, anaponia, refrigerating and brewers' tanks and barrels. In some of these the paint appears to give good results. Nearly all of these paints are simply asphalt or natural bitumen, refued more or less, and a bisulphide of carbon vehicle containing little if any linseed-oil. They evaporate quickly and leave the bitumen coating behind, and probably coat the surface more thoroughly than is possible to apply bitumen hot or in any other manner.

For painting galvanized iron, the bisulphide of carbon appears to be of merit; at least the coatings containing some amount of broulphide either as the principal vehicle or as a drier do not pred as readily as oil paints of similar color and pigments. This favorable point is

soapy conting prevents the oil-paint conting from bonding to the metal, and it dries as a loose skin, peels easily, sometimes before it is dry.

The bisulphide being a solvent of all semi-glutinous substances

The bisulphide being a solvent of all semi-glutinous substances loosens up this soup and incorporates it into the mass of the coating, and the quick evaporation of the bisulphide leaves it there.

There are many instances on record of the disastrous results upon the health of the painters who spread bisulphide-of-curbon mixtures. A noted one is its use with maltha (a mineral bitumen) for the internal and external coatings of a number of miles of steel-riveted water-pipe mains, where the application of this mixture was attended by the disability, insanity, and death of a number of the painters. Its use as an anti-corresive coating for protecting miles of water-pipes was wholly experimental, without a single record on which to base such an application of an untried material, and especially one known to be decidedly inferior and uncertain for minor purposes. Had a gill of this multha point been spread in a room where the Board of Water Commissioners and their Engineering Staff held conneil over the protection of water-pipes from underground corrosion, all the subsequent injury to the painters and expense of application and removal could have been avoided. Another coating was substituted for the malthu. but not before a number of miles of the water-mains had been laid and covered in with no better protection against corrosion than that which could have been had with a coating of boiled skimmed-milk glue,

In the open air bisulphide mixtures can be spread without material danger or discomfort to the painters, but they have not a single element of protective value that warrants their application to any ferric structure of magnitude. They should only be spread on those of minor character, where the corresion or decay is of no material importance, and the question of the cost of the conting and its temporary appearance governs

Frequent analyses of bisulphide paints show about 50 per cent of a low-grade resin, bitumen, and lampblack, for the pigment, with barytes or silica added to give weight. Bisulphide-of-carbon coatings brush out easily and spread over a large area, as the vehicle is very the thermal alone in the country of the armine and the second and the country of the second and the country of the arminest and the country of the second and the country of the country o

Assals new of Carrel darrely for the entire of the entire of the entire of the entire of the entire and the entire of the entire

A summe questionment d'une distre communication de la confliction d'activité de la product de la lacter d'une de la product de la communication de

A freeze de describe est carde en sus a militario de la compensa de la compensa de la compensa de la compensa de la carde en sus de la carde en su

事物 そのなり Aprillador こうかいりゅう コード は、38 。 「いか おまりがり でいまりあい ちゃっか このから #8 機能 一覧 書がある こう 5条子 part はってがける あっとう される あこうが ましまし かっしい ロン 200代表明にはいありない Part しゃ かいいく コート 2004 ロードである それりまいかな いんしゃ であっていた いいん 変名が知るが、10年度のありませる 対象を まる よ おこもい 5月がいる。 それき あいりが ペルルタイル コール・コール・コー

Total and it conton to conton to a product of the state of the second

ess of manufacture, by the action of chlorine on the disulphide of carbon; the reaction being, $CS_2 + 4Cl_2 = CCl_4 + 2SCl_2$. Chlorine, saturated with the vapor of the sulphide of carbon by passing it through the liquid, is passed through a red-hot tube or retort filled with pieces of porcelain, the outlet of the retort being connected to a receiver packed in ice. The condensed yellow mixture of tetrachloride of carbon and chloride of sulphur thereby obtained is slowly added to an excess of potash lye or milk of lime, the mixture being agitated from time to time and afterward distilled. The tetrachloride of carbon passes over, mixed with some of the sulphide of carbon. If too much of the sulphide has been mixed with the chlorine, or if the decomposing heat has not been strong enough, the sulphide of carbon can be removed by leaving it for some time in contact with the potash lye.

No estimated cost of the tetrachloride product is at present.

No estimated cost of the tetrachloride product is at present given, but its field of usefulness in the manufacture of paints and varnishes, also as a special drier, is favorably indicated from the few trials and experiments thus far had with it.

CHAPTER XXI.

JAPAN DIRITES.

JAPAN driers or japans vary greatly in their composition and are very erratic in their action as drying agents. Specimens from the same manufacturer, taken from stock at different times, are widely different in drying qualities, while any attempt to classify the japans of different manufacturers is one of the vexations of the master painters. Probably a good rule for painters to follow in the case of japans is, when one has been found to suit them, to lay aside a sample of it to compare with all future supplies, and to stick to that manufacturer and brand just as long as it comes up to the mark.

The general composition and process of manufacture of japans are: Gum shellae is cooked with lineard oil in a variable kettle until it becomes thick and partakes of the nature of a variable. Litharge and other substances are added to quicken the diving of the resulting product. When the mass has cooked down to a thick substance called a "pill," it is allowed to cool and then thinned down with turpentine. Japan is a light colored brownish vellow liquid of about the consistency of variable. A thin surface of it dives in from 15 to 20 minutes. The care exercised in the manufacturing process and the purity of all the materials used affect sto quality, and are the cause of such erratic results from its use.

The reputation of a japan or various manufacturer counts for much, but it does not always ensure a grant article of the price governs the selection.

Formula for japane are numerous and are trade secrets. The following are representative samples

One gallon cold-pressed old lineard oil 2 peared of 11 C or L. C. gum shellar, 3 pound gold lithstage, 3 pound fortist anchor, 3 pound of

fitharge and red lend, 2 pounds of powdered raw umber. Boil slowly for 2 hours and add by degrees 7½ pounds D. C. gum shellae, and hoil ½ hour longer or until the ingredients are well mixed. Add by degrees 1 pound powdered sulphate of zinc, and when nearly cold, stir in 7 gallons of spirits of turpentine.

The "Bung-hole Drier" formulæ are as numerous as the oil compounders. The following represent a few of the compounds used:

LEAD OILS.

Linseed- or nut-oil	I gallon.	Linseed- or nut-oil 1 goll	on,
Litharge	1 pound.	Litharge 1 pour	nd.
		Sugar of lead k non	md

MANGANESE OILS.

Linseed-oil 1 gallon. Potassium permanganate, 100 grains.	Linseed-oil	
•	of muganese 1 ounce.	

MANGANESE AND LEAD OILS.

MANGANESE A	OND DEAD OIDS,
Linsced-oil,	Linseed-oil,
Linseed-oil, 1 gallon, Permangapate of potash. 4 onnees,	Linsenl-oil
Acetate of lead 4 ounces.	Red lead or lithurge 1 ounce.

See also Boiling Oil, Chapter XXIII. For the effect of different driess upon linseed-oil, see Thorp's experiments, same chapter.

The following is an extract from a Report of "Test on Liquid Driers," read at the Sixth Annual Convention of the Master Painters and Decorators' Association of the United States, held in Detroit, Mich., on Feb. 11, 12, and 13, 4890.

		12											or locatives
						1 11-	aw es) oil es in Min	1100	um ire	l Itri	dyke own es in Min.	l Drie	rest in
						1					·c	100	
1 part	Vzote	drier	(trade-	mark) to	1 part	1	50 35	1	50 50	2	15 20	1	50 20
. 11	**	**	**	20 61	Smirts	12	3.5	2	50	: -1	20	2	20

without showing sediment or separation, called "curdling."

When applied in a thin film to a clean, dry piece of glass placed

in a vertical position, the japun should be dry to the touch in about 2 hours, and should dry hard without becoming brittle in 6 hours.

The so-called concentrated driers are made by heating linsced-oil with lead and manganese salts or oxides in excess, until the product becomes viscous, like a sticking-plaster or birdline. Liquid driers are concentrated driers, thinned out while hot with naphtha or spirits of turpentine. When applied in a thin film to glass and placed in a vertical position, they should be dry to the touch in 2 hours, and harden in about 8 hours. After 48 hours the drier should not rub off in the form of a fine powder when the finger is rubbed briskly over the surface. Liquid driers should mix freely with raw or boiled linsced-oil, turpentine, or benzine in any proportion without showing clots or precipitate after standing 48 hours in the open air.

Inferior liquid driers can be recognized by the odor of benzine when the sample is slightly warmed; by the powdering of the hardened film when rubbed by the finger, and by the rapid evaporation when exposed to the air, with consequent separation of the ingredients.

The quality of a japan depends as much upon its cooking as upon the quality and kind of the materials in its composition. Too high a heat or too long exposure to the heat frequently spoils it.

Gum is added by some manufacturers of japans to harden the oil. This, while causing the japan itself to dry more rapidly, reduces its power to dry an oil paint. Gum is a very uncertain substance in the formula of japan manufacturers. It may mean the spruce end of the schoolgirl, common resin, or the best grade of the fossil resins, over thirty in number, with many varieties in each number.

CHAPTER XXII.

FLAX-PLANT AND LINSEED.

Linserin is the seed product of the *Linum usitatissimum*. This plant is a mitive of India or Eastern Asia, and its cultivation has existed from the earliest ages, distinct evidences of its existence during the Stone Age being preserved to the present day in the rough and worked thix made into bundles, found in the lake dwellings of Switzerland.

It is mentioned in the book of Exodus as one of the products of Egypt in the time of the Pharnols. Among the plagues of Egypt, that of hall destroyed the flax and barley crops, "for the barley was in the ear and the flax was bolled" (Exodus ix, 31).

Pharaoh "arrayed Joseph in vestures of fine linen" (Genesis xli, 42).

Solomon purchased linen yarn in Egypt and Herodotus speaks of the great flux trade of Egypt.

Numerous pictorial representations of the cultivation and preparation of flax are sculptured on the walls and tombs of Thebes, showing the varieties of flax in the red and white flower, the manner of pulling, retting, and hachelling as practised when Jacob dwelt in the land of Goshen; and, except in some minor particulars, or in certain favored locations, are precisely the same as practised at the present day.

The crushing of the seed in a mortar, grinding it on a stone slab by a muller, the pressing out of the oil with stones, the seed-bag, the burning lamp showing that the ancients knew the value of heat to aid in the extraction of the oil, and the painter with his bristle brush and paint-pot is also delineated.

Flax is more extensively and successfully cultivated in Belgium than in any other part of Europe, that raised in East and West Flanders (the Coutrai flax) being the most valuable of the world's crop

flax yielded the following results:

1946 pounds of bolls, which furnished 910 pounds of seed. The
5824 pounds (52 per cent) of flax fibre, lost in steeping 1456 pounds



Fig. 32.—Jerusalem flax-plant blossom. It grows wild in Palestine, covering large areas around Jerusalem. (Blue flower.)

leaving 4368 pounds of retted stalks, and from that 702 pounds of finished fibre were produced. The weight of fibre was equal to about 9 per cent of the dried flax stalk with the seed-bolls, 12 per cent of the bolted straw, and over 16 per cent of the retted straw.

separated in scrutchings, leaving 5.9 tons of finished fibre and 1.47 ton of tow and plackings.

Generally two bushels of linseed are sown per acre, and the yield in finished fibre is from 600 to 800 pounds, the market price of which is about 12 cents per pound. The yield of seed is from 8 to 10 bushels of 52 pounds, and is graded and classified as to quality and condition as closely as any of the grains. The crop is very exhausting to the soil; potash and phosphoric acid are the chief ingredients that the soil requires to produce a good crop of either the fibre or seed. It requires from 400 to 600 pounds of mineral or phosphate fertilizers per acre, beside barnyard and other manures, to keep the soil in condition, and then only two or three crops can be raised in succession, when other crops must be substituted for from 5 to 8 years.

New England formerly raised large quantities of flax for the fibre, but the advent of cotton manufacture soon displaced flax culture, and this, with the exhaustion of the soil and absence of phosphate fertilizers, caused an abandonment of the flax crop in that part of the United States, early in the past century.

America furnishes about one-fourth of the world's supply of linseed-oil. The crop of linseed for the years 1900-1901 was from 16,000,000 to 17,000,000 bushels. The average yield of oil was 18\frac{3}{2} pounds of oil per bashel, or 2,465 gallons of 7\frac{1}{2}-pound oil; equal to 40,000,000 to 42,000,000 gallons. In general, the American crop is comparatively free from the achiteration of the wild unustard and other agrid seeds that render the oil-cake almost valueless for a cattle food. Though, in this respect, it is better than most of the foreign seeds, it is, however, the practice for many seed-crushers to add the screenings from the linseed and grain elevators to their linseed in the crushers, and this not only furnishes a bitter oil-cake but a poorer oil.

The American lineed crop is now chiefly produced by the North-western States, where the rich prairie soil is favorable for a heavy seed crop without much fertilization. The fibre in these States, from its distance to market and the difficulty of preparing it, is of minor importance, and the plant is generally allowed to fully ripen before harvest-

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the world. Flax-growing was begun in Argentina nearly a hundred years ago, but not until about 20 years back was any attempt made

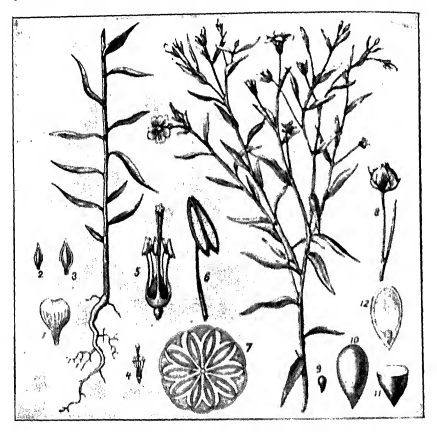


Fig. 33. -- Flax-plant flower, med vessel, and med.

Its flower is blue. Fig. 1 represents a flower leaf or petal; there are five to each flower, which is of a very regular and perfect kind, having five petals, five pistils, five stamens, five sepals. Figs. 2 and 3 are sepals, or cap leaves, to the flower; Figs. 4 and 5 represent the seed-vessel, with its tall stamens and taller pistils; Fig. 6 is a stamen; Fig. 7 is a seed-vessel cut open, showing ten seeds. The stamens fertilize the pistils, the puller falling upon the top of the pistil, or probably carried there by some busy lace. Within each of the pistils (not to

of the most important in the country, and surpasses in magnitude that of any other land. The plant is grown only for the seed, and as soon as the latter is secured the straw is burned. An average of 1000 pounds of seed is raised on an acre, and in some cases the yield is 2000 pounds. export of fluxseed from the four provinces maned amounts to 500,000 tons a year, which is one-half the entire product of the world and

counds 54,200,000 gallons of oil. Not more than 20,000 tons are retained for domestic use, and there appear to be no linseed-oil mills in the country, as all the oil used there is imported. One wonders what the effect upon the markets of the world might be if Argentina should export hisseed-oil and cake justead of raw fluxseed, and could transform the straw into linea thread and cloth instead of burning it.

Ireland, England, Belgium, and Central Europe raise the best flax for fabric purposes, but seeds gathered from these sources being unripe, furnish poor, watery oil.

Russia has a large acreage of flax for seed purposes and furnishes about one-sixth of the world's supply of linseed, the yield being about 8 bushels of 56 pounds to the acre; the flax fibre is of minor importance, being woody and subject to great waste in preparing it for fabric.

Russian seed is exported for seed purposes as well as for oil extrac-In Russia hempseed is sown with the flaxseed, and comprises nearly one-tenth of the seed crop, but as this seed furnishes a siceative oil, it is not an objectionable adulterant, such as the seeds from the rape, colza, mustard, and many other non-drying oil-seeds, called "flax-dodders." The adulteration from these aerid seeds is so great that the waste product in the form of oil-cake, formerly a valuable cattle food, is now so strongly impregnated with the biting taste of these seeds, that eattle refuse to eat it, and it is now used for fuel or fertilizing purposes.

India furnishes about one-eighth of the world's supply of linseed. It is grown as a mixed crop for the seed only. The India flax-plant has been deteriorating for over 200 years, until it is now an inferior shrub from 12 to 16 inches high. The climate is favorable for the oilproducing quality of the seed. The white-flower plant produces about the state of the state of the above an arrest and the state of the sta gathering, the ordinary linseed crop being harvested just before the seed has fully matured, and while it contains more water than if fully riponed.

Rape-seed is sown in large quantities with the lineed. Its yield of seed and oil is very large, and when refined it passes as colza-oil from colesced. These seed-oils are used for burning, lubrication, and in the manufacture of india-rubber articles, because of their non-drying qualities. India is very prolific in oil-bearing seeds; the mustard and many other acrid seeds grow wild, are very rich in oil, and all are freely used to adulterate lineed to an admitted amount of 10 per cent and possibly 15 per cent more.

The quality of the flax, also of the seed, varies quite as much as any crop of grain or vegetables, according to the locality in which they are raised, the soil, weather, and other influences affecting the fibre or oil, and the crop is quite as exhaustive to the soil as wheat or corn.

Samples of linseed grown in various parts of the world and averaged from a collection of ripe seeds weighed from 48 to 52 pounds per bushel, and the yield of oil was quite as variable, viz.:

Gallons of 74-Pound Oil Per 112 Pounds of Spen.

Best Odessa seed	1.5	ter	111	gallous.
Archangel	14		141	**
Good commercial seed	15 5	* *	111	**
East Indian seed	17	**	16 5	**
Sicilian "	iri	3.4	16 5	**
General results by a large crusher for all seeds	 11	* *	17	**

American seed, 521 pounds per bushel, gave 26.55 per cent of oil, or 13.87 pounds.

Linseed in its dry state, as analyzed by Dr. Ure, contains:

Oil.	11	2013 (er cent.
Wax.	11	1 10	** **
Soft resm.	13	\$ 34.54	F2 F4
Resinous coloring unitier	£ 9	2.241	** **
Yellowish coloring matter analogous to tamin.	11	\$12.945	** **
Chin,	11	151	20 20
Vegetable mucilage,	18	1.7	** **
Starch	1	F Ma	
Glutan.	- 1	\$2% Ex1313	** **
Albumin	(FEE	7 1 3 20	** **
Saccharine extractive	100	1 20	** **

Command soluble salts,
Matter insoluble in water but soluble in ether. 4 " " Episperm. 21 per cent.
Soft result and fixed oil. Matter insoluble in water but soluble in other. Matter soluble in water. Water. Water. Eixed oil
Matter insoluble in water but soluble in ether. 18 0 0 0 0 0 0 0 0 0
Analyses by Anderson;*
Albuminous substances. 24,44 per cent Gum and cellulose. 30 73 ° ° ° Oil. 34 00 ° ° ° Ash. 3 33 ° ° ° Water. 7 50 ° ° °
Way's analyses of 33 samples of linserel from various countries;
Nitrogen
Way's analyses, ditto of the oil cake from above samples:
Nitrogen 3 92 to 5 25 per cent. Fnt 6 60 " 15 32 " " " Ash 5 45 " 22 66 " " " Water 6 56 " 10 26 " " " Alluminous substances 25 00 " 36 00 " "
The general composition of all sicentive oils is:
Carbon
Linseed 1 also contains a large quantity of mucilage deposited in the outer layers of cells of the epidermis, which swells up on macerating

boiled in 16 parts of water yields mucilage enough to be drawn out into threads, and forms a dark-colored spongy mass when dry. This crude mucilage contains in addition to the true vegetable mucilage, legumin, albumin, and an organic acid, probably malic acid; also ash constituents, chiefly lime, potash, and iron, partly as phosphates, and partly united in the ash by carbonic acid. Linseed mucilage precipitated by alcohol gives 14 per cent of ash containing 4 per cent of carbonic acid.

Linseed-oil has a specific gravity of 0.928 to 0.953, or 7.743 to 7.952 pounds per United States gallon. The oil from an average quality of ripe seed extracted by various processes contains;

Cold Process.	Hot Process,
Carbon	Carlon 78 11 per cent. Hydrogen
Oxygen 13 85 " "	Oxygen., 10 93 " "
100.00 18 81	1(n) (n) · · · · ·

The earbon-disulphide process gives more oxygen and less earbon. The oil extracted by the naphtha and percolating process does not show any material difference in the quantity of the oil, but is thought to give a quicker drying oil than by the old or cold-drawn process. But, however, it leaves some of the glucerides of the oil in the oil-cake as well as some of the albumin.

The glucerine in the oil in the form of gluceride or other ethers is needed in the change of the fatty acids to form the soap compounds that give the binding quality to the oil. The albuminous substances are organic and are subject to decomposition, and constitute "the drops" or "mucosities" that the boiling process removes, or if the oil is used in its raw state, the driers added are intended to affect them so that they may be oxidized and dried.

Unripe linseed or the seed from flux raised for the fibre (the condition that furnishes a large part of the commercial linseed-oil) contains 5 to 8 per cent of water.

The yield of oil from the different classes (red. blue, or white flower) of linseed varies from 20 to 33 per cent of the weight of the dry seed.

not to exceed 12½ per cent. No. 2. The weight per bushel shall be 50 pounds, the damaged

seed not to exceed 25 per cent.

No. 3. The weight to be not less than 46½ pounds per bushel: the damaged seed, not to be in excess of 20 per cent, is graded "Rejected."

No. 4. No grade. Seed comprises all damp, mouldy, warm seed or those in a heated condition and unfit for temporary storage. All seed that is burnt, smoky, or intermixed with burnt seed is posted as "Burnt or Smoky Flaxscod."

All sales of flaxseed are made upon the basis of pure seed; that is, seed tendered for contract deliveries may carry impure, damaged, or foreign seed matter, but must contain the sale-quantity of pure seed as given, and for such pure seed only shall payment be required.

Linseed yields by the

Cold process of extraction, about 20 per cent of oil. Hot progress " Carlon-disalphide process, 33

Or from 15 to 18 pounds of oil per bushel of No. 1 commercial linseed by the most improved processes of extraction.

The average results in oil of a number of samples of the following substances, extracted by filtration in 100 parts, are as follows:

Almonds. 52 . 41 " Hempweed. 25 87 " Poppy seed. 49 40 28 Grape seeds, 17,95 "

Linseed oil corresponds to the formula (Mulder) Callado, or C_nH₂O₂. It is an hydride of linoleic acid (C₁₈H₂₈O₂). Specific gravity, Linoleic acid is a faint-yellow, limpid oil, insoluble in water. and does not solidify at 18° F., and has both the nature of an oil and a resin. It decreases continually in weight for 90 days, losing from 5 to 8 per cent.

Linoleic acid is peculiar to all sicrative or natural-drying oils, and when fully oxidized by exposure to the air, forms oxylinoleic acid (C₂₂H₃₄O₃) or linoxyn (Mulder).

The linoxyn formed is insoluble in water, dilute acids, alcohol, or ether, and is heavier than water.

Hazura describes linseed-oil as formed of linelic, lineleic, and iso-linelenic acids, and that a high proportion of the two latter acids is characteristic of this oil.

Linseed-oil is composed of drying oil, 80 parts, and non-drying or fatty oils, 20 parts. Of the latter, 8 parts are glycerine ether, the volatile element of the oil, that in the chemical changes among the oil acids by the absorption of oxygen in the process of drying, is absorbed or lost in the change to linoleic acid, with a direct loss in weight of the new oil compound.

The fatty acids in linseed-oil are:

Margarie acid	$C_{ij}\Pi_{ij}O_{ij}$	mercite	gravity	0	810
Palmitic or benic acid. ,					
Olele acid.	$C_{i}H_{i}O_{i}$	**	**	(1	808
Stearie acid	$C_{in}\Pi_{in}G_{j}$	17	**	(1	805

Margarie acid is considered simply as a mixture of stearic and palmitic or benic acid of identical composition $(C_{16}\Pi_{34}\Gamma_{15})$.

The composition and specific gravities of all the fatty oils vary but little, and the influences that affect one affect all.

Oleic other $(C_{20}\Pi_{22}O_2)$, specific gravity, 0.807, associated with the fatty acids, dissolves all solid fats, stearie, pulmitic acids, etc.

The glycerides or glycerine ethers have the characteristics of ethylin (C₈H₁₂O₂); they are the most volatile of the group, and form a component part of the fatty elements in all sicentive oils. The heavy odor recognized when a burning tallow candle is blown out is due to the glycerine ether that comes from the smodlering wick. A heat of 170° F, is adequate to dispel or to cause an absorption of the glycerides into the fatty acids in the oil to form the insoluble scap compound. The albuminous substances in the oil congulate at about 160° F, and in the steamed or hot process of extracting linsered oil, the meal is cooked at 190° to 200° F, to prevent the albumin from flowing out when pressed.

These changes indicate the merits of a low and long-continued heat in the process of boiling oil, instead of the quicker and more

grape, Scotch and silver fir, and spruce. The specific gravities of the whole number (20) range from 0.9202 to 0.9358, varying so little that the hydrometer-test is of little moment to determine their character. The vegetable non-sicentive oils of commercial importance

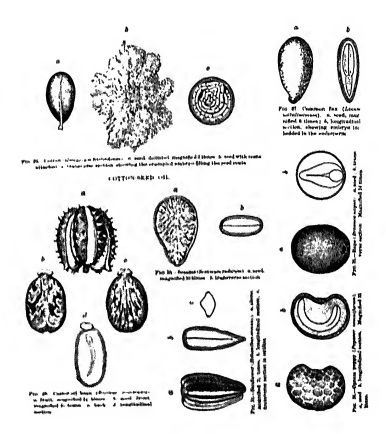


Fig. 34 Oil seeds. (Edlarged.)

are 76 in number, the principal ones being the castor (specific gravity, 0.964), olive, cottonseed (specific gravity, 0.9806, almost identical

burning, and food, are 85 in number.

Vegetable oils, volatile and essential, number 134, or a total of 315 non-drying oils available for the purpose of adulteration. Add 80 animal and fish oils to the above, and it may be conjectured, the source from which a yearly supply from the whole world of 250,000,000 gallons of linseed-oil and other siccative oils can supply a demand for about 350,000,000 gallons of paints, varnishes, japans, and other uses, including the wastes of manufacture.

As before stated, if the flax is raised for the fibre, the seeds do not mature at the same time, hence they furnish a thin watery oil with a greater quantity of the albuminous substances, gum, sugar, and cellulose, called "mucosities," than the 54.44 per cent to 58.50 per cent found in ripe seed. Unripe or mildewed linseed is no more capable of furnishing a good oil than a green apple will make either good cider or good vinegar; an unripe grape, good wine; or grain or corn harvested in the milk will make good bread; and no amount of juggling in the subsequent manipulations of the paint manufacturer can replace the simple operations of nature, or ripen her unripe products or produce them from synthetical compounds.

The following is a comparison of the composition of a few of the principal oil-seeds:

SICCATIVE OILS.

ભા.	Pipovilio Gravily,	('arlsın	Hydrogen	tiaygen,
Linseed Poppy-seed (black) " " (white). Hemp-seed. Walnut	0 9352 0 9270 0 9285 0 9307 0 9288	77 40 76 57 77 20 76 00 176 05 177 15	11 10 11 41 11 31 11 30 11 46	11 30 12 02 11 49 12 70 11 89 11 12

VEGETABLE NON-DRYING OHM.

Cottonseed	0.9155 }	76 40 78 04	11 53 12 04	12 27 0 92
" (summer)	0.9108 (10.174	1 6 174	14 144

RyeWheat			11.79 11.97	11.50 10.84
Barley and oats		յ ե0	11.96 to	12.78 to
Peas and beans	(77.40	11.43 11.30	10.69 12.70
Potatoes and rice Beets, etc			to 11.15	to 11.50

The increase in oxygen in the grain oils is at the expense of the carbon.

MISCELLANEOUS NON-DRYING OILS AND SUBSTANCES.

Sperm-oil	97 10.13
Resin-oil $\left\{ \begin{array}{c} 0.984 \\ \text{to} \\ 0.9910 \end{array} \right\}$ 79.27 10.1	15 10.58
Beef tallow	
Beeswax. 81.60 13.5 Spermaceti. 81.60 12.8	

The vegetable aromatic volatile oils from lemons, oranges, cloves, cinnamon, etc., are of the same chemical composition: Carbon, 88.25; hydrogen, 11.75. When they oxidize, it is at the expense of the carbon element. The diversities in their odor are due to a different arrangement of their atoms.

Poppy-seed furnishes a cold-pressed siccative oil of excellent Being an imported article its price prevents its use except quality. in the finer color paints. The exceptionally good results attendant on the use of French zinc oxide are principally due to the use of this oil. It dries slower than linseed-oil and faster than walnut-oil. not remain sticky so long as linseed-oil, not being so fatty, and takes up less oxygen. It is used for the finer classes of varnishes. It contains 20 per cent of non-drying and 80 per cent of drying oil. Linseedoil clarified by sulphuric acid has generally taken the place of poppyoil. Walnut-oil contains 30 per cent of non-drying and 70 per cent of drying oil. The cold-process oil is light colored with a pleasant smell and taste; after a short exposure to the sunlight it becomes as clear as water. The hot-process oil is characterized by a deep color and unpleasant odor; it has more mucilage in it, and does not dry as well as the cold pressed.

SPECIFIC GRAVITIES OF OILS AND FAIR

Poppy-seed oil 0.4243
Raw linseed cold-drawn oil 0.9299 to 0.932
Boiled linseed-oil
Crude cottonseed-oil
Refined yellow ditto 0.19230
Water-white ditto 0 9288
Menhaden-oil (dark) 0.9292
" (light) 0.9325
Tanner's cod-oil
Porgy-oil 0 9332
Resin, commercial yellow 1 0700
Resin-oil, first run
" third run 0 9887
" " other runs,
Crude Linux petroleum
Lard-oil
Whale- or train-oil 0 925
Sperm-oil p 943
Porpoise-oil
Beef tallow
Mutton tallow

CHAPTER XXIII.

BOILING LINSEED-OIL.

J. Nelson Neil's Experiments.

The study of the composition and properties of the siccative oils has been but little advanced since Mr. J. Nelson Neil was awarded the Isis Gold Medal of the Society of Arts in 1832 for a paper read before that society on "Oil Boiling and Varnish Making" (published in the Transactions of the Society, Vol. XLIX, Part II), which treats so exhaustively upon the modes of manipulation, recipes, and precautions to be used, that the paper is the foundation for all subsequent accounts and modes of manufacturing varnishes. The additions and modifications which have been worked out since that time have not unterially altered the processes of Mr. Neil, either in the betterment of the quality of the varnish or the oil product, but relate more particularly to a shorter time and possibly less cost of manufacture than given by him, which comprised the application of direct fire to the kettles instead of the steam-jacket kettles and devices used by later experimenters and manufacturers.

The substances that act catalytically part with some of their oxygen in the oil, and become to a certain extent deoxidized, and again coming into contact with the air either mechanically blown through the combined mass of oil and driers, or by surface exposure in the settling-tank or barrel, recover their primal condition, and are ready to do the same work over again. It is the necessity for the reoxidation of the driers that causes the general adoption of the air agitation and steam processes now in general use for the boiling of oil and manufacture of varnishes, etc. These driers, without becoming materially altered, induce an alteration in the linseed-oil subjected to their operation. We may imagine this action as similar

the gain in weight after thoroughly dry is from 8 to 10 per cent of the vehicle, and it is heavier than water. Any paint that does not show a marked increase in weight in drying can be set down as one of the hundreds of bastard compounds that masquerade under the guise of paint, with more resin, fish-oil, and hydrocarbon substances in its vehicle than linseed or other siccative oils.

M. E. Cherreul's Experiments.

These experiments of Mr. Neil were followed by the experiments of Mr. M. E. Chevreul, who contributed a paper in 1856 to the Annales de Chime, corroborating Mr. Neil's deductions upon the drying of siccative oils, and by him clearly laid down, viz.:

First. That it is the absorption of exygen by the sicentive oils and the change of the oleic, margaric, and stearic neids of which they are composed, and the chemical combination with each other in the presence of exygen into the lineleic and linelenic acids, that is the cause of their solidification, which term he thinks more clearly defines the action of the oil than drying, which in general may mean evaporation, which is a term adaptable to all liquid bodies, or rather indicates the removal of liquid from all bodies. This definition appears to be apropos to many of the latter-day cheap mixtures called paints; the difficulty experienced with some of which is not to have them dry in a reasonable time, but to have them keep liquid long enough to spread them at all.

Second. That the exidation of the oil is a chemical process and naturally inherent in itself. The action of heat, as in boiling, hastens the drying or resinification of the oil by removing the water and mucosities. That all substances which can be used as driess must be such as are capable of parting with exygen or dissolving in it; and being of themselves exidizable in combination, they in that way increase its absorptive power. There is a class of driess (white copperas, for instance) which act entalytically, while mechanically suspended or in contact with the oil, and increase its exygen absorptive power by their presence, but leave no increase of drying power when withdrawn.

of manganese, it dried in 2 days. When boiled 6 hours with peroxide of manganese that had been used many times before, it required one-half day to dry firm and hard.

Fourth. That manganese driers for exposed paints appear to be less durable than red lead or litharge driers. They harden quicker and become more brittle, from the harder character of the soap they contain, which is further developed in hardness by the heat of the sun.

Fifth. The manganese-drier paints appear to peel more readily than red-lead driers, especially upon ironwork that has but few points or irregularities of the surface to which the paint can adhere. Hence such a paint for an iron surface must remain in a measure softer and more clastic, or else it will be thrown off or peel by the changes in temperature and the expansion and contraction of the metal.

Peroxide of manganese is electro-negative to iron and steel, and is noted for the freedom with which it imparts its oxygen to these metals. It is used in the manufacture of steel as a purge to burn out the imparities in it.

																				•	n()).
Red oxide (of 1	11111	1221	tu:	714	٠.,		ı	, ,		 ,	 	,	84	(15	Ĭu	87	. ()()	per	cent
Oxygen			,			. ,		ï	٠.			 	,	1.1	. 5	×	**	11	. 15	2.6	"
Semputoxide	ní	in	11.				. ,		٠,		 ,	 		1	:	()	••	()	.40	**	"
Alumina.					,				. ,			 		(1		(1)	**	0	. (X)	**	"
Baryin																					
Calcium,																					u
Bilien,	, ,			, ,	, .	, ,				,		 	,	()	. 5	()	**	0	.51	**	"
Water																					"

Sixth. That linseed-oil heated until it lost one-sixth of its weight became thicker, unctuous, and viscid, and dried more readily, forming a tough, crude, turpentine-like mass, scarcely soluble in any other oil (printer's varnish). When linseed-oil is heated to 325° to 375° F, it will take fire and continue to burn without any further application of heat from without, until only tar or charcoal remains.

Nut- and poppy-seed oils also possess this feature, which is sometimes employed as a test of the purity of these oils. If these oils are adulterated to any great extent with fish, resin, or mineral oils,

Four panels of wood were painted on one side with white lead and on the other side with zine oxide, the vehicle being raw linseed oil.

MICHAEL CATE CORE AND ARCHAEL AND AND ARCHAEL AND ARCHAEL AND ARCHAEL AND ARCHAEL AND ARCHAEL AND ARCHAEL ARCH

The results were as follows:

After 24 Hours. After 72 Hours. The white lend nearly set. Set but without adhesion to No. 1. Carbonicthe wood. acid gas. (The zine oxide still fresh. Absolutely fresh. No. 2. Limited air. { The white lend nearly dry. The zine oxide set but not dry. Perfectly dry, The white lend nearly dry, * * . The zine oxide set but not dry, No. 3. Free air. 11 iŧ No. 4. Oxygen gas. Both perfectly dry, 41 18

Professor Vincent's Experiments.

Prof. Chas. W. Vincent's experiments in 1859 were upon the line of boiling oil without the presence of driers, and that a high temperature was not necessary. The temperature used by Mr. Vincent was that due to steam at 40 pounds per square inch, 267° F., used in a steam-jucketed kettle in connection with mechanical agitation by revolving blades and a current of compressed air moderately heated by the act of compression. This process is that in general use at present in the manufacture of lindeum, and it is believed that the Germans practised this method many years preceding 1859 fessor Vincent's conclusions, drawn from experiment, were that air blown through the mass of heated oil is not as important a part of the process as many have assigned to it, and in reality effects nothing toward making the oil a drying one. He boiled lineed oil with air alone, but without driers, for three days consecutively, keeping up a high temperature the whole time, and the resultant boiled oil required precisely the same time to dry as the raw oil from which it was prepared. The body, however, had become so much increased that its consistency was more that of a varnish than an oil.

the air-blast was less greasy and had a greater consistency. Briefly, the surface exposure to the air and the heat secured a sufficient amount of body. The driers produced any required shade of color in the oil and reduced the time of drying from 3 and 4 days, for the raw oil, to 6 hours in the summer and 8 hours in the winter for the boiled. Boiled oil that is subject to long voyages at sea is apt to become fatty and not free working. This is due to the agitation it gets from the motion of the ship while it is under the increase of temperature due to the hold or eargo space in the ship, being, in fact, a long-continued low-temperature and agitation process of boiling. The manufacturer guards against this result by adding to boiled oil for shipment by sea, about one-fourth its volume of raw oil, the oil becoming brighter in consequence of the addition. Professor Vincent's steam-kettle process of boiling gives an oil of a lighter shade than the direct fire or high-temperature process. In both processes, acrylic acid (Callada), a monobasic neid, is produced by the oxidation of acrolein. Acrylic or acroleic acid when purified, is a colorless liquid of a slightly empyremnutic odor, lighter than water and mixable with it in all proportions.

Acrolein (C₃H₄O) is the acid principle produced by the destructive distillation of fatty bodies, resulting from the decomposition of glycerine (C₃H₈O₃). Acrolein is a colorless, limpid, strongly refracting liquid, lighter than water, boils at 126° F.; vapor density, 1.897. Its vapor is so intensely irritating that a few drops diffused in a room are sufficient to render the atmosphere unsupportable. It burns with a clear bright flame, and dissolves in 40 parts of water and readily in other. The solutions at first are neutral, but gradually oxidize and turn acid in contact with the air. Under water it changes into a resinous substance (disacryl-resin) and the water becomes charged with acrylic formic, and acetic acids. The vapor of acrolein passed through a red-hot tube is decomposed with the formation of water and charcoal. Its vapor is highly corrosive to iron bodies.

Acrolein is developed by the decomposition of the nucosities

in the raw linseed-oil under the influence of the boiling heat, and indicates the slower decomposition of these substances in a raw-oil

vehicle for a paint. The slower process of solidification of the raw oil enclosing them, as it were, in a film of the vehicle, to develop later by decomposition into a destructive agent of the paint. The pigments in the paint may delay this decomposing action for a time, but cannot wholly prevent it, and in many cases hasten it. The removal or destructive change of these nucosities is absolutely necessary in baking japans, varnishes, japan driers, and linoleum; and if found detrimental there, it must necessarily follow that they are equally so in a paint oil. The decomposing point in a pure linseed-oil made from thoroughly ripe linseed is nearly 100° F, higher than in an oil made from green or damaged linseed.

Professor Succ's Experiments,

Professor Sace's experiments in brief were, 2500 grains of oil boiled for 10 minutes only, with 30 grains each of litharge and red lead, and weighed after 24 hours' exposure to the atmosphere, the oil had lost only 60 grains. This sample increased 20 per cent in weight after complete resinification. A second sample boiled until there was a loss of 5 per cent in weight of the oil, the product assumed a molasses consistency, and did not resinify after 15 days' exposure to the atmosphere. A third sample boiled to a loss of 12 per cent became a caoutchous-like mass that the atmosphere had no effect upon whatever. It was insoluble in alcohol, ether, chloroform, and bisulphide of earbon; boiling miphtha only dissolved traces of it. The only action which dilute acids had upon it was to extract a small quantity of the oxide of lead due to the driers. Hydrochloric neid dissolved it slowly, while concentrated sulphuric and nitric acid dissolved it rapidly, as they do all vegetable and animal oils and tissucs. This substance was in fact identical with the product obtained by the modern process of boiling oil for the manufacture of linoleum.

Linseed-oil submitted to a dry distillation (without boiling) gave off a white vapor (aerolic) from which was condensed a colorless oil (aerolic acid), having the odor of fresh bread, then expanded and yielded a distillate, a brown empyreumatic product; finally, a mass

of a vehicle to be used for a ferric protective coating. Briefly, the oil is first treated with a dilute sulphuric-acid bath (containing about 30 per cent of sulphuric acid) which is agitated with the oil by the airblast to dehydrate it, but is said to be not strong enough to carbonize it. After standing to allow the oil and acid to separate, the oil is rmi off into the usual steam-jacketed kettle heated to about 267° F., air is blown through the mass, while a solution of manganese linoleate in some hydrocarbon spirit (probably benzine) is added gradually during the process of heating and blowing. Cautionary care is reanired not to add too much of this material. It is the writer's opinion that this caution should extend to the point of not adding any manganese asociated with any hydrocarbon vehicle to the oil, and that prohibition should extend to the use of any sulpharic, nitric, or other caustic acid of any strength to the oil in its preliminary stage. It is almost impossible to clear an oil of either high or low specific gravity, of any acid of whatever strength of solution to which it may be exposed. More or less of the clarifying acid will be held in the oil either free or in combination with the water in the oil, that even a long washing with water, aided by an air-blast agitation, will not remove. Kerosene, napletha, gasolene, etc., are purified by treating with sulphuric acid and their thoroughly washed with water and a long agitation by the air-blast, and are then often found to contain acid enough to perforate the tin cases in which they are shipped. The slight improvement in the color of the boiled oil by this process is a very poor recommendation for its use. The same results are obtainable by the ordinary steam-kettle process, using well-known mineral substances; for itestance, the zine and manganese salts will remove or throw down the inneosities, clear the oil to any desired shade, and cause it to dry promptly, while the water in the oil will be evaporated, naturally by the heat, and the dangers of the sulphur element avoided. Thorp's Experiments with Driers. The action of various mineral and metallic driers upon linseed-

oil in the process of boiling to determine their effect on the color of the oil and the time of drying were made by Mr. Frank H. Thorn, S.B.*

in its favor, the process cannot be recommended for the preparation

and was a very light-colored, cold-pressed, calculate has insect-off, specific gravity, 0.93. The weight of oil under test was in each case the same, 50 c.c. weighing 45.7 grus. The several samples were treated in glass beakers arranged in a sand-bath under temperatures from 200° to 300° F. In general, the temperatures from 230° to 275° F, gave the best results. The time of actual boiling was from 14 to 24 hours, and the percentages of driers varied from less than 1 per cent to 2 per cent by weight of the oil treated. Lithurge furnished an almost colorless oil of firm film, drying in from 6 to 10 hours. Lead carbonate, lead acetate, and lead borate, each furnished slightly colored oils with good films, drying in the order named, in 10, 12, and 20 hours. Red lend, lend chloride, and lend tartrate furnished dark-colored oils of good films, drying in from 20 to 24 hours. Red lead and litharge, 2 per cent of each, also the other lead salts mentioned, with larger percentages than two of each, gave darkcolored oils, all with firm films, drying in about 24 hours. Of the lead driers, litharge gave the best results both in color, film, and drying qualities. Care was necessary in the use of this drier, to not overheat the oil, thus deepening its color. The zine

film, and drying qualities. Care was necessary in the use of this drier, to not overheat the oil, thus deepening its color. The zine salts, the acetate, borate, citrate, oxide, and sulphate, furnished nearly colorless or slightly colored oils with fairly good films, but their time of drying was from 36 to 46 hours. Larger amounts of these driers than 2 per cent shortened the time of drying, darkened the color of the oils, and they did not clarify satisfactority.

The acetate and borate are the best of the zine salt driers, but

the color of the oils, and they did not chrify satisfactorily.

The acetate and borate are the best of the zine salt driers, but none of them act catalytically upon the oil as the lead and manganese driers do, but act mechanically or only while present (like white copperas), to throw down some of the mucosities, but do not east them all out, or set up the combination changes necessary to form the linoleic compounds required in a good drying oil. The manganese salts, viz.: the acetate, borate, sulphate, oxalate, and tartrate, all gave colorless oils, drying with good firm films in from 20 to 36 and 40 hours. The citrate and formate gave slightly darker colored oils, drying in about 24 hours with good firm films. The manganese borate with quantities varying from 1 to 3 per cent of the oil and with

temperatures of 220° to 230° F., gave the best colored and drying oils

some of the samples being tarred. No definite conclusion can be drawn from Mr. Thorp's experiments as to the relation between the quantity of driers dissolved in the oil, and the time of drying of the oil. The action of the several classes of driers, as well as the various members of them, was erratic and the drying result appeared to be governed quite as much by the temperature of the bath, time of boiling, and agitation of the oil during boiling, as by the chemical employed. One per cent by weight of lithurge and the lead driers, and two-tenths of 1 per cent of the manganese salts, were all that were required to give good bright colored oils of good drying qualities with firm films, and not all of these amounts of driers were taken up by the oil, but some were recovered in the residuum.

The sulphate of zine boiled with linseed-oil simply removes the vegetable and mucilinginous substances that impair its drying power; it does not impart any catalytic power to the oil to draw oxygen from the air.

Peroxide of manganese, umber, red lead, and litharge, all dissolve in the oil and impart oxygen to it, and act catalytically to take up more oxygen from the air to renew what they have lost, and in so doing further oxidize the oil.

Thorne and Brin's Oxygen Process.

In this process for oxidizing oil, pure, or nearly pure, oxygen gas in a finely divided stream is poured through the oil at natural temperatures, or moderately heated, if desired. The process occupies from 2 to 7 hours, but a small quantity of the usual driers shortens the time of oxidation. The color and drying qualities of the oil oxidized by this process are excellent. The consumption of oxygen gas varies from 2000 to 4000 cubic feet per ton (250 gallons) of oil, according to the degree of oxidation required.

The principal difficulty in this process is in generating the supply of oxygen gas, which requires a more complicated plant, chain of operations, and intelligence on the part of the workmen than that

The present-day process of boiling oil upon a large scale as practised by most of the crushers of linseed is to dissolve 4 pounds of lead oxide or litharge, or both, in 5 gallons of well-aged and settled linseed-oil at a temperature of about 250° F, for a short time or until all of the oxides are absorbed.

This mixture if allowed to cool will harden into a firm cake of gum (linolate of lead). This product while still hot is mixed with 40 to 50 gallons of linseed-oil heated to the same degree to congulate the albumin, and the mixture allowed to settle. Five hundred or more gallons of this mixture are unde up, and while hot are mixed with a large tank (5000 gallons or so) of raw linseed-oil also heated to about 200° and thoroughly stirred together. This is commercial boiled oil, which varies in character according to the quality of the linseed-oil used in any stage of the process, also in the proportion of the oxide oil to that in the large tank, 4, 6, 8, or 10 to 1, etc.—Lead oxide and a small quantity of manganese oxide make a better drying oil than the red lead alone.

Varnish-makers make this liquid drier for use by local painters, who remove 4 or 5 gallous of ruw oil from the barrel and replace them with the same quantity of the liquid drier, and then roll the barrel around or stir up the mixture with a paddle-stick for their "bung-hole" boiled oil.

Some large users of paint oil think that this nake of "bung-hole" boiled oil is as good as that supplied by the large manufacturers; but this is doubtful, as all of the albuminous substances in the bung-hole oil are retained unchanged, and they are subject to a future decomposition that the 200° of heat in the cooking of the large tank of oil coagulates, and they settle out on standing.

Lead- and manganese-oxide driers made from resin, or resin oil, are marketed on a large scale at 18 to 20 cents per gallon, while a properly made linseed-oil drier cannot be furnished for less than 3 to 4 times that price.

Double-boiled oil means that a double dose of drier and resin-oil has been put through the bung-hole process. The more drier the poorer will be the oil product.

No two manufacturary of hailed all family are it of the

simply a trade-name for an extremely varied composition.

The reputation of the dealer or manufacturer is the best guide. Adulterations by the use of resin-ails are to be especially looked for in all commercial grades of boiled oil.

Boiled oil is a misnomer. Linseed-oil never boils; if it did it would decompose into a permanent gas. The degree of heat applied in the process of so-called boiling by eareful manufacturers is only that necessary to evaporate some of the water natural to the oil. This degree and amount of heat also congulates the mucilage and albuminous substances, so that they are released from the acid oils, and, by their greater weight, are deposited as soon as the oil comes to a state of rest after the heating process. In oil from ripe linsced, if given time to age after crushing, the nucosities and some of the water in its composition (about 5 per cent), that is loosely held in combination, will settle and can be filtered out or drawn off, leaving the oil bright and clear, and with a minimum amount of water to be evaporated in the process of drying as a raw oil. Storage in tanks for 3 or more months still further improves the oil, especially if the tankage is kept at a moderately warm temperature. This fact is taken cognizance of by the varnish manufacturers, who require the best quality of linseed oil for their products, and from the better prices they receive for their wares, can secure the best qualities of

afterward additerated or abused in its application.

As stated before, all driers are injurious to linseed-oil, and the marked inferiority of trade boiled oil to raw linseed-oil is due to the driers used, whether liquid or solid, and not wholly to the removal of the water and mucosities in the boiling process. The salts and oxides of metals that constitute the solid class of driers, that enable the oil to dry promptly, are generally added in great excess of the amount actually necessary for aiding the natural tendency of the oil to dry.

That portion of the driers in excess remains in the dried coating,

oil in the market at a price that the manufacturers of cheap paints cannot compete with. Storage of three or more millions of gallous is carried by some of the best varnish and linseed-oil trade dealers. The natural result of such selection and storage is, that the oil is well aged, clear, and bright, and can be depended upon as a vehicle, unless

CHAPTER NNIV.

DRYING OF LINSEED-OIL.

MULDER's experiments in the drying of sicentive oils determined that there were two periods in the drying of a paint, oil, or varnish coating. The first period occurs in the early months, and is wholly due to the changes in the drying elements of the oil, and while these changes are in progress, the covering is always dry to the touch and remains elastic. This period is of longer duration if the coating is not exposed to the direct rays of the sun. During the period, 100 parts of the oil at ordinary temperatures increase in weight to 111 or 112 parts, but when warmed to 170° F, it loses 4 to 5 parts. In the direct sunlight for all the period of drying, the oil gains about 7 parts.

During the second period the oil becomes hard and firm as a resinous coating, the change being in the non-drying elements of the oil. The increase in weight of the oil is not so great, because the breaking-up of the glycerine element has taken place in the first period. These total changes amount to about a quart of oil in 1000 square feet of painted surface.

The influence of heat in drying a paint or varnish is apparent when it is considered that in the ordinary drying of either to a firm, hard coating, 21 per cent of oxygen has been absorbed from the atmosphere or driers, yet a further exposure to the heat of the sun until the coating becomes hard and resinous ensures a loss of 3 to 5 per cent of this amount. It is to be recommended that this drying and loss be had while the coatings are still clastic, because this loss in substance (due to the changes in the non-drying oil) takes place while the vehicle is soft and clastic enough to adjust itself to the loss in volume.

Oil or varnish dried in the direct hot rays of the sun are not as

peel in strips and cannot be restored; but paint dried in clear cold weather (not frosty weather, that leaves a sweat in and on the conting, as the temperature rises) lasts longer than a sun-dried coating.

It is probable that the cold-dried coating lasts longer for the reason that its change into the hard-soap compound has been dehyed, and being more elastic than the sun-dried, the bond between the pigment, oil, and surface covered is more effectually made. Hence some lead in the from of a drier is, therefore, an advantage in an oil compound or paint, if the pigment does not contain it. The manganese driers are especially unreliable if applied in cold or unfavorable weather.

The percentage gain in weight of a cold-drawn raw linseed-oil, exposed for drying under different conditions, was as follows:

Appleton - Au 1		,				* ****
Daya Exposed.	In Darkmen	l mler Unrhunderl Glass.	Under Huo Glass,	Under Yellow (dam	Under Red Gluss.	Under Green Glass.
10 20	tent tent	0 126 0 236	0.089 0.245	0.012	0.009	0.005
-40	1813 1817	11 258	0 376	0.181	0.082	0.023 0.139
(80) (80)	1113	11 272	0 388 0 357	0,319	$\begin{array}{c} 0.178 \\ 0.338 \end{array}$	0.401
120 150	01% 1138	0 300	0 360	11 142 0 474	$0.376 \\ 0.441$	$0.438 \\ 0.485$
automiconic-d-r a	, 1	1				

After 150 days the gain in weight was the greatest in the following order: green, yellow, red, blue, and uncolored. The application of a dry heat at temperatures of 150° to 170° F, dries a raw oil from 30 to 50 times faster than an open-air exposure under the general conditions of summer weather. Light, heat, and air are all necessary elements to properly dry an oil-paint or varnish coating of any quality. The slow-drying oils gain more weight than the quick-drying.

Raw linseed-oil has a specific gravity of 0.9299 to 0.931, and the same oil boiled, that of 9.411. One hundred parts of drying and non-drying elements in the raw oil in the process of kettle-boiling lose 8 parts of glyceride ether and one part of the carbon and hydrogen from the non-drying oil or fatty acids. This is equal to 9 parts in all, leaving 90 parts that absorb 21 parts of oxygen from the atmosphere when the oil is fully dried, the 100 parts of the raw oil becoming 111 when decided.

on the presence of an excess of drier, or of an unsuitable one. It does not prove adulteration with rosin or rosin oil. It may be due to oxidation of lead linoleate into a liquid turpentine-like body by prolonged exposure to the air. Science indicates no better way of testing a drying varnish than by the finger.

Prof. Max Pettenkofer removed the non-drying acids from freshly dried linsced-oil skins by ether, leaving an elastic caoutchouc-like substance, which by degrees hardened and became brittle. On further exposure to the air it separated easily into thin flakes; in fact, it hardened and cracked like the fossil resins or the coniferm crude gums.

These deductions from a long chain of closely and carefully conducted experiments, not only in the laboratory, but in the application of oil, paint, and varnish coatings, with all classes of pigments, spread over ferric, wood, and mineral surfaces, appear to be ignored in many, if not in most, of the present-day compositions of paints and oils.

The larger amount of uncilage in all unripe or damaged liusced and other vegetable oils, when freshly made, or that have been extracted by the bisulphide of carbon, the hot or steamed seed processes, deteriorates the quality of the oil, and such oil even if it is to be kettle-boiled, is benefited by long standing to allow some of the "mucosities" or "drops" to settle. The amount of these preliminary "foots" is from one to one and one-tenth part in one hundred parts of oil. The greener or poorer the class of seeds from which the oil is made the greater will be the amount of "foots."

CHAPTER XXV.

LINSEED-OIL TESTS FOR ADJULTERATIONS.

THERE are many methods of testing the purity of linseed-oil. The specific gravity test is of small moment, even if not altogether unreliable, as the commercial adulterant oils (over one hundred in number) vary in specific gravity only seven hundredths of a degree Baumé, and no reliable chemical test has been found that is practical for the ordinary analyst.

One difficulty lies in procuring a chemically pure oil to make the comparison with the reactions caused by sulphuric and nitric acids, and caustic-sada treatment for the color changes in the sample tested. The linseed for such an oil must be picked over with the greatest care, selected from fully ripe and full-weight seeds, and pressed in seed bags that have never been used before.

The result of an application to a large number of the most responsible oil-seed-pressing firms, was that not one could furnish a sample with less than 5 per cent of other seed-oil in it, and most of the best commercial brands contained over 12 per cent.

Approximation chemical tests are numerous, but in all cases a sample of known quality of linseed-oil should be used for comperison, even if it is not a chemically pure oil. Abbets and other refractometers are used to test the purity of linseed-oil, but their use is too complicated for any one but an experienced chemist.

All the fatty oils change color when brought into contact with strong sulphuric acid. If a drop be added to 8 or 10 drops of an oil placed on a glass plate resting on white paper, the following colors are immediately produced. Olive-oil turns a deep vellow, gradually becoming green. Sesame-oil, a bright red. Colza oil, a greenish-blue aureola. Poppy-seed oil, a pale yellow with a dingy gray look. Hempseed-oil, a distinct organic dinger. Justice by the production of the production.

IV, p. 183.

Mineral oil or petroleum in any form cannot be added to linseedoil to exceed 5 per cent without affecting its drying; and 10 per
cent prevents its drying other than as a thin skin impervious to the
air, and the oil remaining green beneath is liable to blister or peel on
exposure to simlight. It does not hond to the pigment or surface
conted.

If a tin plate conted with a mixture of a vegetable and a mineral oil he viewed at different angles in strong sunlight, the mineral oil can be detected by the iridescent or metallic play of color, which petroleum imparts to all vegetable oils. So characteristic is this, that a little experience will enable a painter to readily detect mineral oil when present even in a small quantity. This feature is in some degree disguised or palliated by treating the mineral mixed oil with caustic lime, chalk, etc., and adding an excess of strong free driers, which may suppress the iridescent base; but if the sample is ignited, the marked pungent odor of burning petroleum vapor is readily recognized.

Cottonseed oil, a semi sicentive oil, was formerly mixed in large quantities with linseed oil. Its specific gravity, color, taste, and odor are almost identical with linseed oil. It requires a large amount of driers either compounded with it by heat or as free driers to enable it to dry. Its tendency at the best is to harden and crack the overlying coatings; also to crack in cold weather, from its non-clastic condition due to the driers used. Crude cottonseed-oil treated with strong ammonia mixed with 3 parts of water, gives an opaque brown color. Refined cottonseed oil, similarly treated, gives a brownish or dull opaque vellow. Pure linseed-oil similarly treated for comparison, gives a bright, but semi transparent yellow. Mixtures of both oils give an intermediatory color, that a little experience will enable a painter to determine approximately the amount of the mixture.

A quick test by cold is to place a sample of the cottonsed-oil, mixture on a piece of glass alongside of a known sample of linseed-oil, and place the glass in a refrigerator or on a piece of ice. In a short time the impure sample will become discernibly thicker than the linseed-oil

use to adulterate the latter has sensibly diminished. There is, however, a combination of the damaged seed oils from both crops, for which there is always a demand to furnish a mogret hisseed-oil at a cut-rate price.

Resin oils are freely used to adulterate lineard oil, even by firms whose business reputations should warrant more honest wares. Resin mixes readily with lineard-oil, and whether its grade be light or heavy, it cannot be detected by its specific gravity. For a quick test of its presence, shake up a spoonful of the sample with 5 times the quantity of strong alcohol; pour off the alcohol and add to it a clear solution of sugar of lead; a cloudy precipitate shows the presence of resin.

Resin-oil can also be detected by the remarkably manscons after-

taste produced by it when touched by the tongue. The odor of the oil is not recognizable in the sample unless ignited, when it becomes decidedly different in odor from ignited lineard oil. If the cork of the sample bottle squeaks when it is twisted around in the neck of the bottle, no further test is necessary to denote its presence.

Lineard-oil adulterated with resin-oil of any grade is readily de-

tected by passing a current of chlorine gas through the sample oil, which is rapidly blackened if any appreciable amount of resin is present.

Resin-oil is especially to be looked for in boiled oil. It never hardens completely, and makes the conting "tacky." If any considerable amount of resin is present in a lineard oil that has received an excess of driers to harden the coating, the conting will be brittle and crumble easily on a short exposure to the atmosphere, particularly in summer weather, or by exposure to heat.

Menhaden and porgy fish-oils are used freely to adulterate linears-oil, especially in many of the mixed paints and pastes. The price of these oils is only about one-half that of a poor Insection. Fish-oil in the twentieth century, used as an adulterant of lineard oil, comprises almost anything from a whale to a mossbanker, with the oil from dead animals frequently added to help out the aboundation. The fish-oils dry slowly, but surely, if fortified by strong, free driers. Fish-oils used for a tin-roofing paint will stick longer than a lineard-oil paint, as they do not dry so hard. They are more affected by cold than lineard-oil, and whatever paint conting is suread over one with a

when inoderately neared.

Place a sample of a known quality of linseed-oil and one of the oils to be tested, in separate test tubes, cork, and then heat together in a hot water-bath; if the suspected oil gives off an odor of aerokin (oxidized glyceride) similar to that of the smoldering wick of a tallow candle, fish-oil of some grade and amount is present.

One of the most reliable tests of the purity of linseed-oil, and one that does not have to be felt for as in the preceding tests, and is equally available, is, viz.: Add to 100 parts of the oil by weight, one-half of one per cent by weight each, of htlange and red lead well stirred together. Heat in any convenient vessel, and in any manner, until an immersed thermometer reaches 480° to 500° F. A feather from a feather duster or chicken's wing will answer instead of a thermometer. feather when dipped in the lot oil curls up with a crackling sound, it indicates an approximate temperature. A small current of air from a bellows or other source should be blown through the oil as it is being gradually heated. A small glass tube and a piece of druggist's rubber tubing are readily available for this part of the apparatus. Small samples are taken out from time to time and cooked on an iron plate. As soon as the samples appear stringy when cold, allow the oil to rood, stirring it constantly during the cooling. If the oil is solid and farm where cold the sample is of good quality. A poor oil will be sticky and more or less third, and of bad other,

Animal oils can be detected by their odor when the sample is heated, also by the addition of nitrie or sulphuric neid, either of which gives an intense-red color to fish oils. The adulteration of mineral oils is readily discovered by the process of suponification, when these substances rise to the surface. The suponification number, or the number of milligrams of Is HO required to suponify linseed-oil, is 190.2 to 192.7. This number for many adulterated oils is as low as 180. The judice number of linseed-oil is 158.7 to 159.78; that of fatty acids, 159.85.

It is frequently necessary to clarify linseed-oil. The following are a few of the methods:

Heat the oil slowly up to 300° C, (570° E) either by itself or with the addition of from 1 to 5 parts of either caustic line, carbonate of Mulder recommends the clearing of turbid huserd oil by washing it with its own volume of warm water containing some common salt.

Sulphuric acid is used for clarifying Insection, particularly adulterations of it. Its action is to carbonize the fibrine elements of the fish-oils and the inneilinginous substances in the vegetable oils, and to deposit them in the so-called "foots" or "drops," Its action is injurious to linsect-oil in general, for it removes by carbonization a part of the fatty acids, or non-drying elements, and all of the glyceride others, the latter being essential to the changes necessary to form a firm, lard cont from the oil when dry. Acid-treated oils are liable to dry on the exterior only, and never become hard or firm, Acid-treated oils require long, careful, and repeated washings with warm water in the form of an air-blown spray through the body of the oil in a deep tank to climinate the acid, which is seldom thoroughly done. The acid, besides delaying the drying, will attack the metallic-base pigments afterward associated with it in the paint.

Graphite and carbon pigments are less affected by sulphuric acid in the oil than any other class of pigments.

Graphite paint-skins detached from the metallic plates on which they had been spread and dried, when immersed in 5 per cent solutions of sulphuric acid, lost in weight from 1.5 to 1.7 per cent, but were not affected in lastro, strength, or clasticity.

This result indicates their superior qualities for heavy contings for roofing paint, and in other locations where any sulphur element can reach them, whether from the vehicle or the atmosphere.

The results of Mulder's experiments with sulphuric acid treated oils to ascertain their qualities were that they did not begin to dry materially under 8 days. At the end of 3 months the samples had gained 15 per cent in weight, but lost 3 per cent of this when heated to 150° F, for a short time. Also a strong heat from the sun for a number of summer days produced the same effect.

Pure linserd-oil, not treated, gained 10 per cent in the same period and lost 2.5 to 3 per cent on heating it. The acid did not affect the drying elements in the oil, only the non-drying ones, as noted before.

The colors of the soid-treated oils were not contained offered by

ing character, if ground in a hot mill will develop this feature more fully.

Sulphuric acid in the oil or pigment appears to take kindly to wooden surfaces as a priming coat; at least the disintegrating effect of the acid is not so marked as upon metal. All sulphuric-acid-treated oils have a tendency to increase the galvanic action in all paints made from them, that are spread on wooden surfaces. When used on metallic bodies electrolysis is increased.

The following table * gives the weight in grains of sulphur in a gallan, of a number of the oils when burned by means of a wick floating in the oil, and condensing in a sulphur apparatus the vapors of combustion, the same way as sulphur is determined in coal-gas:

Name of Oil.	Grains of Sulphy Per Gallon.
Linseed-oil (La Plata)	trace.
Cottonseed-oil	trace.
Olive-oil	none.
Groundnut-oil.	none.
Sperm-oil, ordinary.	2.3
" Inttle mae	3.1
Coronnut-oil.	0.1
Nentsfoot-oil	3.7
Col oil	4.7
Burn all therefore	5.8
Rape-oil (Jamba).	11.3
intia, otionii) * * * * * * * * * * * * * * * * * *	14.2
" " ordinary brown	17.4
" I brown, relined with sulphuric acid	16.8
" " " " fullers earth	10.0
" " (Ravinistr's)	10 1
Russian mineral oil, erude, o sus.	20.5
" " Intring	10.3
American mineral oil, water white	8.1
" " Imrining	10.1
tt tt mefanter	16.3
to the safety	14.0
Scotch mineral oil, used for making gas	49.8

does not separate to any great degree, unless it stands for many weeks, and then being at the bottom of the can or package, escapes notice.

The covering or spreading power as well as the coloring power of such mixtures are of the poorest character; at least an extra cont and often two are necessary to present any sort of a creditable appearance. They dry solely by evaporation through the outer skin of the paint, leaving it porous, and where moisture can escape, the same element containing other atmospheric gases can enter and they are more destructive than the moisture they replace. All water oil mixtures dry flat and lifeless. The use of alkalies and strong metallic salt driets to form a better emulsion does not materially change their forced mechanical association, and if they cannot evaporate and escape outwardly, they go inward and condense on the covered surface, form blisters, and peeling results.

It is almost impossible to spread a water-oil paint in the cold without heating it. If spread and not dry, a cold night, not even approximating a frost, will ensure a blistering and pecling the next day. Brushing the coating over with turpentine or bearine will not prevent or correct this action, which will take place regardless of the nature of the pigment. A good linseed-oil paint spread on a cold day (not freezing weather) will "take" and dry if a little extra effort on the painter's part is made to brush it out well and by using a little more turpentine for the drier than that used in warmer weather. But all painting for external exposures should be suspended in cold weather, especially on all ferric structures, unless under cover and in a warm room, where the painted surface should remain until the paint has at least "set" firmly, or until it has dried enough to bear handling without feeling "tacky."

Many paint chemists deem that 2 per cent of water added to linsced-oil in excess of that natural to it, whether made from ripe or unripe linseed, is not detrinental to a paint. To enable the oil made from unripe seed to earry the extra water, it is rendered slightly alkaline, generally by adding lime-water, which forms with the oil vehicle a calcium soup that thickens the paint, so that it never settles hard, and is easily stirred up, consequently, does not dry hard.

A number of tests for free water in linseed-oil are: Heating the oil

refrigerator, and note the difference in its spreading. Heat a piece of iron to a bright cherry red and phinge it into the oil or paint. If there is much snapping, it indicates the presence of free water in the mixture. In ordinarily pure oil or good paint, a thick, heavy smoke without explosion or snapping will follow the withdrawal of the glowing test-iron.

free water and swell up. Cool the paint or oil for a few hours in a

CHAPTER XXVI.

SUBSTITUTES FOR LINSEED-OIL.

Many so-called substitutes for linseed-oil have been presented to the public in the past, and at present they are municipally the hundreds in the Patent-office records, and outside in the formulae of the compaunders of paints. All substitute oils are as uncertain and indefinite in character as the pigments assembled with them.

Generally, a low grade of linseed-oil is the base for the vehicle, to which one or more of the score of thax-dodders or buffuns, resin, fish, mineral, india-rubber, and soap-fat oils are added. These are mixed with benzine for the volatile, also with manganese or other strong metallic-salt driers, put through the bung-hole. No heat is employed in their manufacture, and some of them are dangerous to sell, or spread even when cold.

Purchasing agents and master painters (except in a few cases) have not the laboratory, chemical apparatus, technical knowledge, or time to analyze them to detect the frand. The result of their use comes with the lapse of a very short time, when the scraper, burning torch, and a new conting are the only remedies for the evils of crazing, peeling, or decomposition due to galvanic action. No amount of sophistry can change the fact that the use of so-called substitute oils has in nearly every case been detrimental to the paint and the covered surface.

Probably 80 per cent of all the oils, paints, and varnishes nonunfactured in the world is applied to structures of minor importance, which are destroyed by causes other than corrosion. These contings are quite as much for looks as for physical condition, but the other 20 per cent is used on the most important and costly engineering structures of our time. These require protection from corrosion from the hour the materials leave the rolling-mill, forge, and foundry, antil they are in the faither attention as a transfer of the continuous structures.

made, viz.:

"Lucol" * is a paint oil, a synthetical compound, made by a secret process from materials that necessarily are a part of the manufacturer's secret."

What it is as a chemical compound, or as a manufactured paint oil, concerns the proprietors of it. What its claims are for superiority over linseed-oil, concerns the users. As set forth by its manufacturers, its characteristic features in comparison with linseed-oil are:

"1. Lucol is more durable than linsced-oil, which dries by absorption and oxidation and only to a small degree by evaporation. The reverse is the case with lucol, which dries principally by evaporation, hence the condition of the atmosphere to which it is exposed while drying must be considered.

"2. Lucol sets quicker than linseed-oil, which is the result of evaporation instead of the oxidation of its elements; the final drying to a bone-hard condition requires many months. The retention of its elasticity no doubt accounts for its durability.

"3. Lucol sets sooner and dries quicker than linseed-oil, hence is less adhesive for dust, and does not wash off if rained on, as is the case with other vehicles.

"4. Lucol gives a purer white with white lend than linseed-oil.

"5. Lucol preserves the original tint of the pigment longer. More lucol is used in a coating than when linseed-oil is used. The gloss is less at first than with lineerd-oil, but it is soon ahead in this respect. The oil is the life of the paint.

"6. Lucal can be flatted with a much smaller proportion of turpentine than with lineced-oil."

Other advantages are set forth, but all are more adaptable for the use of lucid on surfaces other than ferric, and have been answered elsewhere in this work.

The Lucal Company says: "We extract the olein, which is carefully refined by a special and partly secret process, and by the use of chemicals it is converted into a brilliant, transparent, lemon-colored oil. It contains no regetable, mineral, or fish oil, resin or resin-oil, varnish-gams, benzine, or other powerful solvent driers.

an unpleasant odor while being spread and in drying, wholly unlike linseed-oil, and should be flowed on similar to a varnish, instead of being brushed out like a linseed-oil coating.

"Lucol, in the form of a paint, resists alkaline substances, senair, sen-water, and covers galvanized-iron surfaces without peeling. Lucol weighs 7½ pounds per gallon, and is placed on the market on its me its as a *synthetical manufactured oil*, wholly anlike any other substitute compound heretofore offered as a paint-oil."

How well are the above chins founded? "We extract the olein," etc. This at once destroys the chain that lucol is a synthetical compound. It is only an oil made with a vegetable, an animal, or a fatty acid base.

All of the solid fats and oils are derived from two sources. The marine animal oils are obtained from the cold-blooded fish, like the cod, menhaden, etc., and the hot-blooded, like the seal, sperm, and right whale, etc. The terrestrial animal oils are lard, near's foot, horsebone, tallow (oleic neid), etc. Both classes may be considered as mixed glycorides of oleic acid $(C_{18}\Pi_{32}O_2)$, stence neid $(C_{48}\Pi_{30}O_2)$, and palmitic or benic acid $(C_{18}\Pi_{32}O_2)$; the first preponderating in the oils and the two last, especially the stearie (stea ine), in the fats.

Oleic acid has a specific gravity of 0.808 at 65° F, and is the liquid acid obtained by the saponification of non-drying oils and liquid fats, which contain a different glyceride than the drying oils. The proportion of olein differs according to the nature of the fat four which it is obtained.

Chevreul prepared it by boiling human fat, laid, goose-fat, beef, and mutton suct, filtering the solution and allowing it to stand for 24 hours, then concentrating it a little by evaporation, adding water to separate the olein, and separating the liquid from the solid anotter by pressure. Olein thus obtained does not solidify at 32° F.

Olein is also prepared from olive-oil and other glycerides containing it by pouring upon the fat a cold strong solution of caustic soda, which suponifies the solid fats but not the olein. It is also obtained from olive and almond-oils by treating them with cold alcohol and evaporating the solution.

Pure olein is a colorless oil void of taste and smell, insoluble in

olein will have an odor distinctive of the class of fats from which it is obtained.

The narrine oils all have the repulsive fishy odor in various degrees, sperm-oil being the hardest to locate. The terrestrial animal oils have the peculiar sourish odor of cooking fats. The vegetable oils have a sweetish odor. A little practice with a heated sample will enable the most of them to be recognized.

Olein is also extracted from the organic acids in soap-stock or the fats left in the by products in the refining of cottonseed-oil. The fatty acids in the "foots" are distilled with superheated steam; when the distillate cools and solidities, the dein is extracted by pressure. The process is analogous to the production of commercial cottonseed-oil and lard stearins used in the preparation of butterine, lard surrogates, and candles. There are about 250,000 gallons of olein available in the cotton crop of the United States, if all the foots were used for the extraction of olem and none used for the namufacture of cheap soaps.

There is no amount of animal oils or fatty refuse available for numufacturing into olein that can in any material way affect the broad field covered by buseed-oil as a vehicle for paints.

Chemistry has not arrived at that stage of development where the assembling of the chemical clements of fatty substances in their known proportions will produce an oil or fat. All such substances must have a mitural base for the foundation for the chain of operations and reactions necessary to change their nature.

Liked, as a paint vehicle, therefore, is not a synthetical companied, but a manufactured paint oil. Its endorsement by unster painters when used on passenger cars or other works which are covered by contings of feesal gum variash, or upon ferric bodies which are thoroughly covered with rust, is no evidence of its resistance to corresion. A few successful applications of it under favorable conditions will not counterbalance the failures. Generally, no record of the failures of substitutes for hissest-oil is available for the public, who are as much interested in knowing what not to use, as what vehicle is the best for a paint

of drying thoroughly in about 6 to 8 nours. Figure its ground in the oil furnish excellent paints, that do not remain soft and sticky below the surface, like contings prepared from linescal oil.

Chinese wood-oil is favorably employed as a floor oil or paint on account of its hardness; also in the manufacture of an oilcloth-like goods, which, when dried in hot air, excel the ordinary oilcloth or water-proof products by their extraordinary elasticity. The odor of

the oil is very peculiar, resembling land, and remains in the coating for months, and even for years. This land odor remains in the lacquers made from the oil; hence for that use, also for their and other interior uses, it is necessary to remove it. Disguising the small by the use of a volatile oil does not answer the purpose, because the odor reappears after the evaporation of it. Among the remedies resorted to are; Agitation with a dilute solution of permanganate of potassium; a filtered solution of chloride of lime, filtration through animal charcoal; mixing with potato flour, also storing it for a long time after filtration, after the process of Bang and Ruffin. It is also possible to obtain a tolerable freedom from the odor by the use of a blower passing air heated to not exceeding 50° C, through the oil, for 6 to 8 hours, when it loses perceptibly in odor and can be used for lacquers or floor-work. For outside exposures it is not necessary to attach

of mucilage and albumin. In this state the oil becomes waxy at low temperatures, and organic salts analogous to the stearates settle out. Wood-oil is beiled for a short time in a like manner to linseed-oil with a small percentage of red lead or litharge, else it will always remain

Wood-oil in its raw state dries opaque, probably due to the presence

much value to the dedorization.

with a small percentage of red lead or litharge, else it will always remain opaque. This for paints is immaterial, but boiling is necessary to give a greater drying quality.

In boiling the oil, whether with lead or manganic compounds, a

temperature of 200° C, must not be exceeded, otherwise, in the use of borate of manganese, a thickening cusues, followed in a short time by complete gelatination and waste of the oil. Heats approximating 160° C, and in any case not over 180° C, should only be used, and for but a short time, when the oil should be removed from the fire and the siccatives stirred in. This imparts to the oil a drying quality and

from the wood-cil, and the elasticity from the linseed-cil.

The important drying quality renders wood-cil useful in the manufacture of fatty lacquers. It cannot be employed for spirit lacquers, as it is insoluble in alcohol.

The wood-oil of China and Japan is obtained from the seeds of the tung-oil or varnish tree (Aleurites cordata, Elacococca vernicia). Another variety, the Aleurita triloba, furnishes an oil of less drying power, and is used to adulterate the oil from the former.

About 266,700,000 pounds of the oil are annually shipped from Hankow to other parts of China, and for export. The Canton woodoil is said to be better and purer than the Hankow, and is about 10 per cent higher in price. The cost of these oils in England, where their use is firmly established, is from 4 to 4) cents per pound.

De Negri and Shurlati report that the fruit of the tree from which the oil is extracted contains 53.35 per cent of oil, 42 per cent being recoverable.

The cold pressed oil is of a pale yellow color, is tasteless, and has a smell like easter oil. The hot pressed oil is a medium brown in color, with a taste and smell like hog fat.

The drying power of the oil is superior to that of linseed-oil, the cold-pressed drying better than the hot pressed.

Its specific gravity is 0.936 to 0.941. Suponification value, 156.6-172. Iodine value, 159-161 (de Negri and Shurlati). The oil is soluble in cold absolute alcohol and mixes readily with linsced-oil. When heated with litharge it turns darker in color and evolves a

slight smell of aerolein. When thinly spread on glass in a closed room it dries to a whitish thin resembling milky or frosted glass. A heavier conting exposed in the open air dries in about 6 hours. The oil after

heating or boiling by itself develops the whitish film, but when boiled with litharge is as clear and bright as any oil varnish.

Very thick layers of the dried oil can be scraped off as a tough mass quite uniform throughout. It has an exceptionally small adherence to glass.

The bulsain known as wood oil or gurjan bulsain, from the Dipterocurpus turbinatus, Guerta, should not be confounded with the tung-

Euphorbium.

This substance is in its experimental stage in the United States as a vehicle for anti-corrosive and anti-fonling paints. Attention was directed to its anti-corrosive qualities as a natural varnish and its probable utility as a vehicle for paints about the year 1870, from the discovery that the axes, machetes, and other tools used to ent down the thickets of the euphorbia spurge, to clear the way for a surveying expedition in Natal, became coated with a strong glutinous inice that adhered so firmly to them as to be with difficulty removed. The tools conted with it did not rust in fresh water, and hilge water had but little effect upon it. When articles were control with it and immersed in the sea, no barnucles or marine life would adhere to it. Its effect upon insect life appeared to be equally repulsive, and timber conted with it resisted the ravages of the Teredo navalis. It resists heat and cold better than linseed-oil and varnish vehicles, while ammoniacal and chemical vapors do not cause blistering, scaling, or other injurious action.

Euphorbium juice has a strong affinity for iron and steel, and when applied in its crude state as it exades from the shrub, has no injurious effect upon metals, wood, or other substances used for engineering or common building purposes. When prepared for a paint, the juice undergoes several special processes and becomes a clear gummy vehicle of a medium-brown color, that receives the usual color pigments much as linseed-oil does; retaining, however, its own protective properties unimpaired.

Euphorbium, prepared for a vehicle, appears to maintain its properties in all climates, and does not apparently deteriorate with age. It is perfectly clastic, tenacions, and when dry, can be drawn out to a thin thread. It adheres firmly to polished steel, tin plate, zinc contings, sheet lead, and spelter. Earth acids appear to have little effect upon it, as pipes conted with it and buried for a number of years show little injury.

Euphorbium juice has a bitter, biting taste, is very acrid and irritating to human flesh, corroding and alcerating the body wherever it is applied. The sores resemble those from nitric acid, and are bord to head. In this and negative that there are a total acid, and are

The euphorbium of commerce is imported in casks, and is a gummy, resinous substance in the torm of drops of an irregular size resembling gum arabic. The drops contain vegetable matter twigs, flowers, thorus, etc., that collect on the gum as it exides from the tree and are dried in; though many of the tears are hollow and without refuse in them. The natural color of the tears is a cloudy pale yellow externally, but of a lighter color internally. The tears break easily in the fingers, but are difficult to pulverize. The principal part of the process to prepare the crude gum for use as a varnish or a vehicle for paint is to free it from the vegetable matter. It is partially dissolved by water and almost entirely by alcohol, ether, and oil of turpentine. Its composition by analysis is, viz

Vients whilds in other	. 26	95	wr	cent,
re a madullik " "	. 11	25	٠,,	11
Emphation, the presching principle	. 31	141	**	**
Chantelman	1	tn	**	**
Malic mals	1	503		**
Giggs maltin	2118	Iti	**	4.6
Animoma aduble matters	1	Зu	**	**
	1183	tici	**	**

It has no neid reaction but an extremely burning taste,

The interese acridity v due to the result soluble in the ether, which melts at 107 6 to 109 4 1. The result is obtailed in ether melts between 246.4° and 248.4°. Fuplication is a crystallizable substance, fusing at 154.4° 1, and soluble in ether, because, etc., but not in hot water.

Emphortum direct cashly without the use of metallic salts or solvent driefs. Ox gall and other kindred substances appear to be the liest driefs for it, when any are required. The crude resin is the product of the Lapharhan or—the genus is trainetons. There are about 600 species, many of which are natives of hearly every country in the temperate zones, and are commonly known as spurgeworts.

The explications sparse, or I reconfere is a shirilder and herbiscome succulent frequently covered with those and having stalks from 3 to 6 and sometimes 10 feet in height, and grows in almost imponeThe euphorbium gum from Natal is considered to be inferior to the Morocco product. North Africa is capable of producing cuphor-

bium in sufficient amounts to supply almost any demand for it,

The gum is called in Arabia "Darkmows," and is known in the

Eastern markets as "Farfinm."

In India there are 116 species of the Ameurdacear referred to 23

genera, in addition to the E. resinifera and some other varieties found in Arabia.

The *E. dracunculoides* in India (*jy chee*) yields 25 per cent of a clear oil of a yellowish or greenish-yellow color from the dry husked seed. The oil is more limpled than lineared oil, does not become ropy from age, and is used for a burning and drying oil.

The E. lathyris is raised on the edges of the fields in France, Germany, and Switzerland. It contains 40 per cent of a fluid oil of a siccative nature.

The *E. neriifolia* grows wild in Burma, Baluchistan, the Mulay Islands, etc. It yields a gum of a gutta-percha nature on boiling the stems and twigs of the shrub.

The E. Royleana is a large fleshy shrub common on the dry rocky hillsides of the Himalayas, growing at an elevation of 6000 feet. The sap of this plant yields a superior gutta-percha.

The Pisticia Lentiscus vields the resin mustic,

The Melanorshwa usitatissima yields the black varnish of Burma.

The Indian Holigarna longifolia also yields a varnish.

The Indian Odina Wodier is covered with its brown gum, which streaks down the stem and ultimately turns black.

The *E. pulcherrima* is an originated shrub grown in Mexico that yields a milky sap which fardens into a black gum, and can be boiled down to a sort of gutta-percha. Guatemala and other countries near the torrid zone also have a large number of trees that furnish the natural varnishes, though no attempt has been made to bring them into commercial importance.

The P. terebinthus is a tree growing along the shores of the Mediterranean Sea. It furnishes the cyprus turpentine.

The Japanese lacquer-tree, or the urushi-naki, is known in China

wood, ivy, etc. The American dogwood was formerly thought to be of the same species, but is now placed in another of the same order.

In Japan * the lacquer-tree grows to the height of about 30 feet, and at the age of 40 years is about 40 inches in diameter. It reaches its greatest perfection in the yield and quality of the lac or varnish at the age of 18 years.

The crude lac, called ki-urusha, is collected at any time between the months of April and October by making a number of horizontal incisions in the bark of the tree ma manner similar to the "boxing" practised to gather the sap of the long-leaf pines or turpentine-tree. The tree is backed in this manner for from 60 to 80 days, or until it dies, when it is cut down, the bark and sap-wood removed and steeped in hot water, which extracts the last remnant of the lac, about half a pint, which forms the poorest quality of he, known as "roiro-urushi," or black varnish. The tree seldom survives the first season's backing, at whatever age it is done.

The varnish-tree is probably native to China, but it is also found native in Japan, and is cultivated all over Nippon and in several districts of Kinshia and Shikoku, and there are extensive plantations in the valley of Tadamigawa and Northern Echigo. A temperate climate scenes to best suit the growth of the tree, as it reaches its greatest perfection on the main island north of latitude 36°. It is cultivated in Northern Hondo, between 37° and 39°. It may be of interest in considering the question of habitat to note that the Rhus vernicifera, mentioned by Mr. J. J. Rein, are growing in Germany at Frankfort on-Main and at Strasburg. They endured the hard winter of 1879 80, when the temperature reached 27° C. In Japan the lowest temperature in Northern Nonshin is ~ 12° C.

The lac is parified by straining it through cotton cloth, evaporating the water by exposure to the sun or by a gentle heat. Sometimes water is added to the crude lac, and they are ground together on a paint slab, and then the water is evaporated. Various coloring matters are added to the purified lac by grinding, as is usual in the manufacture of oil paints. Black lacquer other than that furnished

gall from pigs and oxen, to give body to the lac. The purest lac is from the first run of the sap after tapping. It is called *mashyi urushi*, and is bleached in shallow vessels laid in the smulight. The other principal grades are the *henki-urushi*, the unbleached *j shimi-urushi*, and the *roiro-urushi*, or black varnish.

There are about 20 different grades and qualities of these lacquers in the Japanese market, of which the above are the principal ones. They vary in color from a light brown to a deep jetty black.

Lacquer is thinned only by heating. The addition of water thickens the lacinto a jelly. Lacquered objects are always hardened in a humid atmosphere, such as a room with wet cloths hung on the walls, or containing a spray or vessels of water.

All varieties of varnish-trees are propagated by the seeds and

cuttings. The seeds are gathered in October and sown early in the spring, make 10 to 12 inches' growth the first season, and in 10 years are 9 to 10 feet high and from 2 to 3 inches in diameter. In a favorable soil the annual height-growth during the first 6 years is from 20 to 30 inches, and diminishes afterward to from 10 to 20 inches. Plants from root cuttings grow more rapidly than seedlings, but the latter make hardier and longer lived trees. The trees after the first 5 or 6 years require very little care, and are generally tapped at my period after the tenth year of their growth, though sometimes it is done when only 4 or 5 years old.

The climate and soil of at least one-third of the United States

are as favorable for the growth of lacquer-trees as those of Japan or China. Their cultivation requires no more care than that given to the sugar-maple or the Encalyptus. Specimens of the trees are growing in the grounds of the Department of Agriculture at Washington, D. C.

Plants which are largely cultivated in Europe have been confused with the Japanese Rhus remicifera. They are, however, a different variety of the tree. The Ailanthus glandulosa Desf., in France called Vernis de Japan, is also of the variety-bearing species.

The poison-sumach, Rhus verneuta, common in the Eastern United States, yields a sap that furnishes a black, historis, durable varnish, very similar to that derived from the Japanese tree.

yield as white or pure a lacquer as those in China or Japan.

A species of varuish-tree that grows in India was thought to be the veritable Anacarde, but it is entirely different from the Japanese "urusi" variety.

No attempt to cultivate any of the varnish-trees on a commercial scale has been made in either Europe or America. The manufacture of lacquer and lacquer ware is one of the most important industries of China and Japan. It seems natural that if the largest users of varnish in the world depend almost solely upon these natural products, their cultivation in America is well worth trying.

In China the Rhus vernicifera, or variish-tree, is called Ch'i-shu (Tsi-chou), also Li tschi. It grows wild in the province of Fingo and on the island of Tricom, and is cultivated in the mountains of Hupeh and Seechwan, but the best varieties are found in the province of Januatto, where it is cultivated extensively. It is probable that the Mingpo and Fouchow varnishes, as well as the Hupeh, are from the Rhus vernicifera—A varnish-tree growing in South China differs from the above variety, but is not well known at present.

In China the Rhus vernucifera grows about 15 to 20 feet in height, seldom reaching one foot in diameter, and has but few branches. The bark is white, knotty, and peels readily. The wood is fragile, resembling the willow; the pith is very abundant. Its leaves have a mild taste, and when rubbed on paper, dye it a dull black. The flowers are greenish vellow, and have an odor resembling orange-blossoms. The fruit is of the size and shape of chick-pea, and at its maturity is very hard and of a duty color. The seed furnishes an oil and wax which are extensively used.

From the berries of the Rhus vernicifera, Rhus succedaba, and other related Chinese and Japanese species, a vegetable tallow is extracted and used for candles. The wax is exported in large quantities to Great Aritain and the United States for an adulterant or substitute for beesway.

The general composition of cende lac is lac acid (a resinous acid, soluble in other), 60 to 80 per cent, a gum 3 to 6 per cent, a nitrogeneous substance resembling albumin 17 to 3.5 per cent of a volatile acid, and water, which are driven off in the preparation of refined

inhaled or in contact with the flesh, produces what are known as varnish boils, accompanied by an intolerable itching and burning sensation, similar to that produced by the poison ryy. They are difficult to heal, and resemble the effect of nitric acid on the flesh.

The Chinese and Japanese use because as a varieth or vehicle

for colors on all kinds of household utensils, also for the inside and outside coatings on their buildings. Lacquer as a vehicle can be used for all colors except a pure white and some of the lighter shades of other colors. It is applied to wood, porcelain, and metals, and forms a hard resinous surface, highly lustrous, practically insoluble in boiling water, alcoholic liquids, alkaline and acid solutions, nuless in a highly concentrated form. Applications of lacquer to the underwater surfaces of a number of Japanese war vessels, for both anti-rorrosive and anti-fouling contings have been very successful. The coatings, after a sen-duty of four years of the vessels to which they were applied, showed no signs of either feating or corresion. Applications of other anti-fouling paints of all characters over lacquer coatings were failures, the urushic acid of the lacquer attacking the metallic base of the foreign anti-fouling paint, resulting practically in the destruction of both, The best results for under-water marine work with lacquer is had

The best results for under-water marine work with heeper is had when the first conting is a heavy one and almost pure lacquer. The succeeding coats can be thinner in body and contain either a pigment or some inert substance to give body. Mica, graphite, lampblack, etc., have been used experimentally with success for these secondary coatings.

CHAPTER XXVII.

DECAY OF PAINT.

Rustr proceeds solely from the action of an acidulated moisture upon a bright or clean iron surface, and is probably only a point at its inaugural. The affinity of the iron for the oxygen in the acidulated moisture of the air or water in the oil, or from other sources, is greater than its bond with the hydrogen as water (H2O), the decomposition ensuing releases the hydrogen, which is sixteen times the volume of oxygen united with the iron to form hydrated Fe_sO_s or red rust. The hydrogen, from its light specific gravity, in its effort to escape into the air pushes up the overlying paint coating, increases the area of the affected part, cracks the conting in its exit, moisture enters again, and corrosion is master of that location. The rust which has thus been formed is hygroscopic and carries 24 per cent of moisture as it forms. This moisture never dries out under any atmospheric bent conditions, but is ever ready for a chemical decomposition; the hydrated red rust, being nearly two times the volume of the iron from which it is formed, adds its efforts to the free hydrogen to push up the coating and form a blister and crack in the coating, How energetic this mechanical action due to corrosion is, may be observed on the ordinary east-iron hand railings for fences and outside steps of New York City and other city houses, which in hundreds of instances are split for more or less of their length. Cast-iron water or gas pipes, with bell and spigot joints, are frequently unade with rust joints. They almost invariably burst the bells by the swelling of the iron cement used to make the joint,

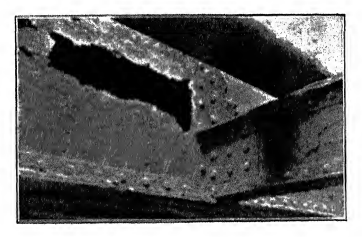
The cut, Fig. 35, shows a section of a well-known railway viaduct, the iron construction having been painted over mill-scale, or in the condition the naterial left the rolling-mill and workshop. It had received the usual treatment given by contracting engineers to remove the mill-scale preliminary to painting.

were pineed in position.

The corrosion existing in Brunell's* tubular iron bridge over the St. Lawrence River at Montreal, Canada, had proceeded to so great an extent as to require the removal of the whole structure, it being impossible to repair it. The efforts to replace the cross floor-beams supporting the rad stringers resulted in loosening every rivet in the neighborhood of the repairs. Pitting around the heads of the rivets had proceeded so far and deep that it was impossible to cut them out without loosening every configuous rivet.

This bridge had been kept well printed with iron-oxide and some experimental paints applied contrafter cont. These contings whenever removed, or that fell off during the periodical element, or attempts to repair and paint the structure, showed the several coatings of millscale, paint, and new rust formations as plainly arranged as the leaves of a book.

Fig. 37 shows a similar state or corresion.



In. 37 Corrosion of steel girder Washington Street railway bridge, Boston,

In 1879, Sir Nathan Barnales stated as the result of his observictions of chips' metal in the English naval stations, that when the mill-scale was left upon the plates, angles, and other parts of the 11. It is see a see a see a see a second residual as an est restur situal In 1887, Mr. Rialton Dixon gave before the Institute of Naval Architects his experience as to a vessel built entirely of steel some eight years before, which was found to be greatly corroded in the bunkers and water-ballast chambers near the engine room and boilers. Some of the anglesirons had entirely disappeared, and the tic plates were eaten away in holes. This action could be traced directly to the presence of milt-scale, and whether the surfaces were conted with paint or cement or not, the corrosion was always present upon those plates and angles that had toill scale upon them, and was absent in those free from it. The presence of the paint or other coating retarded corrosion only in a minor degree by preventing moisture from reaching the metal covered by the mill scale.

In 1882 Mr, Farquianson, on behalt of the English Board of Admiralty, conducted a mumber of very exhaustive experiments at the different naval stations to test the action of mill-scale on ships' metal. The result was to establish beyond dispute that, first, no pitting occurred on mild steel when freed from mill scale; second that the loss in weight from corresion of clean mild steel and clean from did not differ much; third, that the action of trail scale is considerable and continuous, and equal to a similar amount of capper in its corresive action on metal covered by it. Since these experiments the Admiralty have never wavered in their practice of having all of the ships' metal pickhed to remove the mill scale, whether it is to be covered by paint or cement, or to be galvanized.

Destructive Agents of Points

Pure water is a greater destructive element to an oil conting than solutions of sal annuouse, chloride of magnesium, common salt, or natural sea-water, if free from sewage, all of which are agents of destruction. The decay of a paint is hastened by mechanical action if the water, either fresh or salt, or the other solutions, are in motion. Ordinary commercial oil coatings are destroyed by diluted nominatic and nitric acids, alkaline liquors, ammonia, sulphide of ammonium, soda, caustic alkalies, and alkaline solutions of coal nelnes, clinkets, einders, soot, etc. Diluted sulphoric acid days not materially affect

A conting of paint appears to be a very simple thing, as it is, when applied to a louise or barn and both are left to their fate, but when applied to an important engineering structure, with all the vicissitudes of service in the extremes of heat and cold, sunshine and storm. atmospheric and other gases from natural or manufacturing sources. from corrosive liquids and solids, it is a different matter, and requires more engineering experience to select, more chemical knowledge to compound, and more technical details to get the right thing in the right place at the right time, in the right manner, and in the right amount than the general run of master painters do or can give to the subject. If the influences to which a coating of paint is to be subjected are known, it can generally be determined in advance whether it will be durable. For instance, zine white or oxide (ZnO, specific gravity, 5.42) amplied as an external coating absorbs carbonic acid from the air and some moisture, changing to a carbonate of zinc (ZaCO_a, specific gravity, 144). During this change there is an increase in volume from 119, as an oxide, to 28.1 as a carbonate. This change from an axide to a carbonate is a chemical one, and occurs during the process of drying, but the change in the volume of the two substances exerts a mechanical action also in the atoms of the pigment, not only to disrupt theto and leave them loose and easily carried away by the wind, rain, etc., but cracks and loosens the oil vehicle in which the pigment is embedded no well no its bond to whatever surface it covers But if the zine oxide conting is applied in a closed room, though the gir contains the same amount of carbonic acid, or even more than the external air, the exide close and change to a carbonate, as the necessary moisture is lacking; hence sine oxide for internal contings is durable, but for outside contings is perishable.

Red lead (Ph₂O₂) specific gravity, 9.07) remains unchanged under ordinary atmospheric conditions, but if the air contains hydric sulphide, as it does in many manufacturing establishments and towns, to a notable extent, it will be an inexorable chemical law change the oxide to a sulphide of lead (PhS, specific gravity, 7.13), and this chemical change (usually denoted by the blackening or discoloration of the east) will also be accompanied by an increase in volume cination of the copperas or roasting of the ore tachererofore noted). is another instance in which an inexcende chemical chance in one of the pigment's loose substances is accompanied by a change in i.s. specific gravity, its corresponding change in volume, and a torchanical action to reinforce the chemical action due to the raw oil vehicle loaded with its charge of driers, whose function is to either decorepose or consume by a slow combustion the "macristics" in the oil while attempting to dry. All these instances are similar in effect to what would occur in the plastered wall of a building if the mortar used in it, when partially dry, should begin to not one in volume to the amounts as given above. Other instances could be cited, but these show that the pigments of the conting can be so chosen as to proclude the destruction by them of the coating, but that it is almost impossible to guard the vehicle from the injurious influences inforent in the composition of the pigment, that is chanced an elementer, after its application, by chemical laws. Hence the absolute necessity that an order for a protective paint should include the conditions it is to be subjected to,

In addition to the preceding remarks upon iron oxide, graphite, and other paints, and the several tests given in detail of a rew of the many paint compounds, it may be noted, viz:

All pigments * can be grouped into three class a possisting to their affinity for linseed-oil.

First. Those that form chemical combinations called some and are generally the most durable. They consist of lead, zinc, and iron bases, of which red lead combines with the oil to the createst extent; next, the pure carbonate white lead made by the "tild Durch Process," followed by zinc oxide and iron oxide, Turkey umber, vellow ochre; also, faintly, the chromates of lead, chromospicen, and chrone yellow.

Second. Pigments of this class, being neutral, have no chemical affinity for the oil; they need large amounts of driers, either combined with and carried by the oil, or as free driets. They include all blacks, graphites, slates, slags, vermilions, Prussian, Paris, and Chinese blues, terra de sienna, Vandyke brown, Paris green, verdigris, ultramarine, carmine, and madder lakes. The last seven are trans-

parent colors, and are better adapted for varnish mixtures and glazing.

Third. Pigments of this class act destructively to linseed-oil, They have an acid base (mostly tin salt, hydrochloride of tin, and redwood dye) which forms, with the albuminous and gelatinous matters in the oil, a jelly like compound that does not work well under the brush nor harden sufficiently, and can be used in a varnish for glazing only. Among the most troublesome are the lower grades of so-called carmines, unadder lakes, rose-pinks, etc., which contain more or less acidulous dyes, forming with linseed-oil a soft paint, that dries on the surface only and can be peeled off like the skin of ripe fruit.

"Catalysis" is a term introduced by Berzelius, and by him applied to the changes that sugar solutions undergo in the process of fermentation, and now used to denote the changes that certain substances, by their mere presence, effect in other bodies without themselves undergoing may apparent change. Catalytic action is a potential agent in the decay of point contings, and manifestly has not received the attention from paint chemists and compounders that its marked nction on the life of a contain warrants. The present efficiency of the inemplement gualisht is wholly due to catalytic action between the substances that compace the mantle when excited by the comhustion of the gas. In the development of this light all of the rare mineral existes and metals and the exists of the baser metals, chromining alumina, codestt, manganese, nickel, and iron, when associated with thoria in the mantle, have been found to act as entalytic agents to carry, condence, or absorb oxygen, that increases the flame temperature of the mantle and consequently increases the light. This flame temperature in some cases reaches the point where volatilization of some of the baset metals and oxides custos. Clinconl. powdered glass, percelain, their spar, crystallized quartz, pumice-stone, and other kindred substances are also found to act as entalytic agents in combaction, but do not develop so high a flatur temperature in the mantle as the either entertaines above moted.

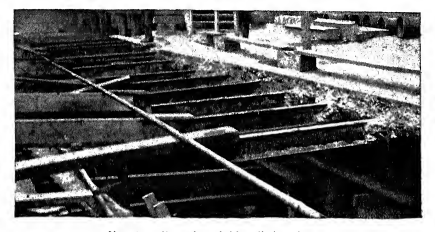
Combustion of any substance may be quick and attended by a

catalytic to some other substance, either in its own class as a so-called inert mineral pigment, or in the chemical class of oxides having a lead. zine, iron, or other metallic base. Individually, they may be apparently unaffected by long exposure to the air while in their loose state or in packages or bulk; but when mixed together, they take up moisture or oxygen to a greater or less degree, either by absorption in mass or by condensation upon their surfaces, and catalytic action ensues. The oil vehicle and driers are catalytic of thenselves, and when mixed with the pigments act more energetically as carriers of oxygen even when the coating is apparently dry. In all pigments and vehicles, the one that is the most refractory, or that is the most resistant to exidation in whatever form the exygen may be presented. is the one that nets the part of the thoria in the gaslight mantle, becoming the negative or non-consumable substance, that, though excited to a greater activity by the presence of the other substances in the paint compound, retains its resistance to a change the longest at the expense of the other associated substances. Thus far, lampblack and graphite, in their subdivided form as pigments, appear to be the only substances not subject to catalytic action, or if it is present it is so weak that the life of the conting is not uniterially affected from this cause.*

Caustic Action of Mortar upon Paint,

An examination (1901) of iron floor-beams taken out after an exposure of about forty years showed that the beams originally were particularly well painted and laid in a location where only the dry warm atmosphere of a residence reached them. The paint coatings had been thoroughly destroyed by the caustic action of the lime mortar used to turn the brick arches in which the beams were embedded. Corrosion was well established in every inch of their surface. Had any moisture, as in the case of the Times building, reached them, their condition would have been fully as bad.

The iron beams supporting the sidewalks laid about forty years ago in New York City, that were removed for the Rapid Transit Tunnel work, invariably show deep corrosion from the destruction promoter of it, if moisture or air can reach the surface so covered.



Fra. 38. - Corrosion of sidewalk iron beams.

Hydraulic, also quickline mortar, only prevent corrosion so far as they are free from mill-scale and continuously dry to exclude the air. The paint coating when burned by the caustic action of mortar or cement, adds no material period to the life of the iron and except for appearance and protection during construction, might be left off. (See Chapter XV.)

The modern hollow tiles used for floor arches and building partitions with their advantages over brickwork, do not remove the cause of the corrosion of any iron that they may be in contact with.

Gypsum, while not caustic, is hydrometic, and the continual presence of moisture is fatal to ferric bodies; besides, it is not always free from caustic substances developed in the calcination of it.

The following eement for the levelling, bedding, and in contact with metal work, is recommended. The cement hardens like stone, is impervious to water, and can be applied by a trowel from a mortar-board, over walls or to lay brick wherever mortar can be used. It is made from marble dust (from marble sawing or pulverizing mills) mixed, viz.:

cement porous. When the mastic is to be used, for every 100 parts of such mixture, 7 parts of linseed-oil are required to bring it to a good trowel paste. The oil can be either raw or boiled, according to the time of drying required. The surfaces to which it is to be applied should be dry, clean, and preferably coated with linseed-oil or a good carbon paint, before the application of the cement.

A refined bitumen coating applied to the bright metal, hot, has proven to be the best of coatings for ironwork laid in cement, mortar, or concrete, to correct the caustic action of them.

The metal work of the movable dam at Lake Wennibicskish, Minn., constructed in 1899–1900, was cleaned bright by the sand-blast and then painted three coats of Edward Smith's Co.'s Durable Paint, applied one week apart, each coating being thoroughly dry before the application of the next. Observation of the paint in 1901 showed that the coatings had been completely killed and absorbed wherever the painted metal was embedded in the concrete. The metal was as clean as before painting, with a slight discoloration of the surface of the concrete from the paint absorbed. The metal exposed, however, did not show the same tendency to rust quickly, as before the application of the paint, on the short exposure before again being put in place. The surfaces not in contact with concrete were in good condition.

CHAPTER XXVIII.

SAND-BLAST AND PICKLING PROCESSES.

THE sand-blast is the most satisfactory and simplest method of cleaning all surfaces for painting, whether at the shops or in situ.

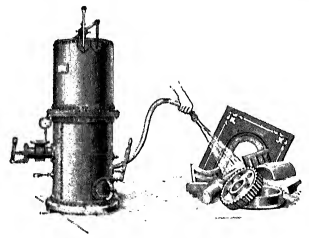


Fig. 39. Sand-blast apparatus.

The invention of the sand-blast is due to General Benj. Tilghman, and was patented October 18, 1870, No. 108,408, but has since expired. There are some patents for sand-blast apparatus of subsequent date, issued to other parties for improvements relative to portability, clogging of the sand in the case, etc., still in effect.

Fig. 40 shows a portable sand-blast apparatus used by Mr. Geo. W. Lilly, C.E., for cleaning railway viaduets in the city of Columbus, Ohio.

The principal features of the sand-blast consist in the use of compressed air at a pressure of from 15 to 25 pounds per square inch, discharged through one or more chilled-iron or hardened-steel

per hour for each nozzle, which discharges about 120 cubic feet of free air per hour, or about 1 cubic foot of sand to 1000 enbic feet of free air per hour. The nozzles wear rapidly and require frequent

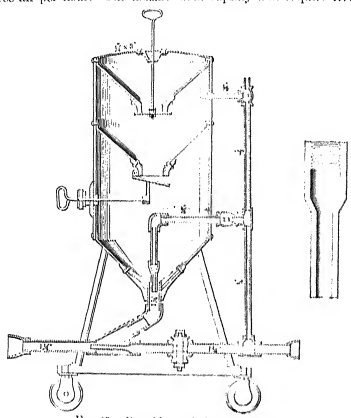


Fig. 40. Portable sand-blast machine.

renewal, but they are of small or minor expense. The 2 or 24 inch diameter armor-clad rubber hose that conducts the compressed air from the air-receiver to the place of work being soft and elastic, is comparatively little affected by the current of sand and air, unless the air is hot; hence methods to cool the air before it reaches the leading hose are necessary. Four such nozzles that gradually wear to 4-inch diameter and then discharge 200 to 240 cubic feet of air

paint coatings are generally thick and somewhat softened by the water, about $\frac{8}{10}$ square foot of metal is cleaned per minute, or 48 to 60 square feet per hour. This costs about 3 cents per square foot of surface, as the waste of sand is greater and the work cannot be done so advantageously in a dry dock as in a shop.

On the New York Elevated Railway Station at 155th Street an average of 80 square feet per hour was maintained for a number of months in removing a hard coating of old paint and rust to the bright iron. The loss of time in changing nozzles and shifting scaffolds was about one hour per day per nozzle. The labor account was one man to hold and direct each nozzle; one man to attend to two sand-boxes, and one man to clean up and supply sand for the four nozzles or seven men per corps the men shifting their scaffolds without other aid. The four-nozzle plant for bridge or viaduet cleaning will clean about 2500 square feet of surface per eight-hour day at an expense of about \$20 for all itens, except the man and coal for the compressor, or 8 cents per square foot, which would be modified by the amount of cleaning for each structure.

Removal at the shop of mill-scale and dirt is done at the rate of 4½ to 5 square feet of surface per minute, or 270 to 300 square feet per hour per nozzle, or about ¼ cent per square foot of surface. With an organized corps and plant, the cost of cleaning surfaces need not exceed ¼ cent per square foot of surface, large or small, or about one-third the labor-cost of the painter on a first-class coat of paint, and requires about the same degree of skilled labor as painting.

The metal-work of the movable dam at Lake Wennibioskish, Minn., erected during 1900, was cleaned bright by an extemporized sand-blast. A hoisting-engine run backward furnished the compressed air, an old steam-boiler was used for an air-receiver, gaspipe for nozzles, and garden hose for leaders, etc.

Mr.W. C.Weeks, C.E.,* reports "that four laborers and one engineer in charge of the apparatus cost for labor \$9.22 and \$2.50 for fuel. On general surfaces, 40 square feet per hour were cleaned, using two nozzles,

In a number of United States navy-yards, with well-equipped, permanent, and fairly perfect sand-blast plants, the cost of cleaning averages ½ cent per square foot of surface. This for plates ¼ inch thick is 98 cents; for ¼-inch plates, \$1.95 to \$2.00 per ton. To sand-blast 7-inch 1-beams weighing 17.5 pounds per foot costs \$1.35 per ton; 12-inch 1-beams, weight 50 pounds per foot, cost 80 cents per ton.

The average cost for cleaning plate-girder bridges in situ is probably \$1.00 per ton of metal. For trass and lattice-iron bridges the cost of sand-blasting ranges from \$1.00 to \$1.75 per ton.

The United States Army Engineer Corps cleaned 50,000 square feet of steel lock-gates and other metal on the Muscle Shoals Canal during 1898-99 from a temporary floating plant. The cost of all items, was 3 cents per square foot, and the new coat of paint cost 2.88 cents per square foot.

Mr. Geo. W. Lilly, C.E., reports the cleaning by sand-blast of a number of railway bridges and viaduets in the city of Columbus, Ohio. The work was done under exceptionally adverse circumstances, but indicated that 8 cents per square foot covered all the expenses. For cleaning a viaduet over the Little Miami-Railway, containing 25,000 square feet of surface in a contined location where the cleaning was interrupted by the train-service that frequently amounted to one-fifth of the working hours, the cost of the work, including flagman, sand and drying, compressed air, and all other expenses, including the labor, was 3.04 cents per square foot. On the best days, in a favorable location uninterrupted by the trains, 1227 square feet were cleaned per day at a cost of 1.23 cents per square foot. On a plate girder containing 3727 square feet, the cost was 2.37 cents per square foot.

The pressure of air ranged from 25 to 38 pounds, averaging 33 pounds per square inch. The nozzles were rapidly. They were of ½-inch extra heavy wrought-iron gas-pipe, about 2 feet long and lasted from 3 to 5 hours each.

The expense of handling structural metal at the shops after machining preparatory to sand-blasting it, ranges from 40 cents to \$1.50 per ton according to the weight and character of the pieces and facili-

A recognition of the fact that structural steel is a perishable numerial, requiring thorough protection from corrosion during all the stages of its manufacture and use, should be required of every engineer, and the subject should form an important part of his education. There is no part of structural engineering needing a more thorough reform in both spirit and practice than this one.

The apparent indifference regarding the future fate of steel material, after it is in location, is probably due to the mistaken economy of the engineer corps and the proprietors, on account of the added cost of properly cleaning the metal. If cleaning is necessary, as engineers and all admit, it should be specified in the contract, properly done, inspected by a competent person, and paid for like any of the other processes, and the penalty for its non-fulfilment be as strictly enforced as for a badly driven rivet or poorly machined or fitting part. The above deficiencies are readily detected and can be corrected, but the poorly cleaned surface escapes notice and is readily put out of evidence by the lumdy paint-pot.

It is the imperative duty of the engineer in charge of structural work to require his inspector to perform his duty at all times so that a radical change shall be had from the present practice of eleming and painting ferric metal, the corrosion of which is now too much in evidence.

At a late meeting of an engineering society, the protection of

ferrie structures from corrosion was under consideration by oral discussion and correspondence. The cleaning of the surface of steel to the absolutely clean metal by some method, preliminary to the immediate painting of it under cover, was unqualifiedly endorsed. And yet within the limits of a ten-mile circle from the engineers' meeting there were many thousands of tons of ferric structural material in process of crection by the engineers represented at the meeting, and scarcely a ton of this material had received any other cleaning than that from a putty knife or a whisk-broom. The quality of the applied paint in many cases was as deficient as the cleaning. So much for theory versus practice.

Pickling the metal instead of sand-blasting it is more practised by European than American engineers, especially for structural work. are recommended. They are equally as safe to handle and should be applied hot if possible; the latter quality is best for removing the scale.

In foreign navy-yards, 9 to 10 per cent hot solutions are used, the metal remaining in the bath five or more hours, according to the quality of the scale. This requires a large pickling plant and has no other advantages. When appearance or test shows the scale is loosened, the metal is removed and well washed by a copious and strong jet of water under 75 pounds or more pressure.

Soaking the metal in baths of still or light running water does not thoroughly remove the acid. The still-water bath is the cause of the failure of tin-plate.

Pickled metal is liable to become conted with a tough, gunnny substance, quite difficult to remove, except by the friction from a strong jet of water. Arsenic in the sulphuric acid made from pyrites also adds to the gunnny deposit precipitated on the metal. Acid free from arsenic should be specified for the pickle. The gunnny deposit prevents the paint from bonding to the metal, rendering it liable to peel.

After the metal has been washed by the jet of hot or cold water it should be immersed in a bath of hot lime-water and be left in it long enough to reach the temperature of the bath, in order to neutralize any of the acid not removed by the water jet. It is then removed and dried, preferably in an oven. The conting of lime left upon the metal can be easily brashed off, leaving the metal clean and bright, which will show evidences of rust in an hour if not painted immediately.

Murintic acid is sometimes used in place of sulphuric acid for the pickle. It is not as effective as sulphuric acid, it costs more, and the gummy coating formed by the pickle is more difficult to remove, requiring a hot alkaline or caustic-soda bath, instead of lime, to remove it. A solution of sulphate of zinc is effective for the removal of this gummy coating.

Obviously a pickling plant requires a larger stage for an equal

cleaning biths, and ovens, will be from 50 cents to \$2.00 per ton, or rather more than is required for a sand-blast, as the several pieces, though of the same weight and character, have to be moved more frequently.

Steels high in earbon, or cast-iron articles, are difficult to pickle, as a film of graphitic curbon forms on the surface of the metal, which mixes with the gummy deposit from the acid bath, and requires considerable labor and care to remove.

When the sand-blast or pickling process is not available, mill-scale, rust, and old paint coatings are removed from works in situ, by the gasoline burning torch, followed closely by the scraper and wire brush.

The cost of the burning process is so closely connected with the painter's labor as to be difficult of separation, but a quart-burning torch will burn $3\frac{1}{2}$ hours, and one man can saturate rust-spots and burn off from 80 to 100 square feet of surface per hour, at a cost of $\frac{10}{10}$ cent per square foot of surface, leaving it ready for the painter.

A modern parlor, or sleeping-ear, 65 to 70 feet long, requires three gallons of gasoline to burn off the *outside* paint coating, and about four days of time, for one man to use the torch, followed by two men two days each, to sandpaper rendy for the painter, or a total cost of 45 cents for the gasoline, and \$15.00 for the labor, or $\frac{8}{10}$ to 1 cent per square foot of surface.

Carc is required in the use of gasoline, either for the torch, or for saturating the old paint, as explosions and serious burning of the workmen, and fires, are frequent. Insurance companies forbid the use of either the torch or fluid for the removal of paint or rust in any building covered by their policies.

Any material that can be inclosed in a chamber or iron easing and subjected to the action of a bath of low-pressure steam for 20 to 25 minutes will have the old coatings softened, when they can be easily scraped off. This is to be followed by a thorough washing with soap and water, and rinsing. A pair of locomotive driving-wheels required 30 minutes to scrape and wash after steaming. The

pounds of melted laundry or soft sonp. The two mixtures added together, when cold, should be of the consistency of putty. It is applied by a trowel or stiff 4-inch flat brush in successive coats about $\frac{1}{8}$ to $\frac{1}{4}$ inch thick. Care must be used in mixing the lime. A stirring-paddle should be left in the tub to form a vent to prevent the enustic mixture from blowing out.

The cost to remove the paint from a pair of locomotive drivingwheels by this mixture is 65 cents for material and 15 hours of labor at \$2.25, or a total of \$2.90. Careful washing with hot water to remove all traces of the caustic-soda paste is required, as for all strong alkaline mixtures.

Wooden surfaces treated with caustic-soda compounds to remove paint or varnish are injured by the raising of the grain of the wood, which cannot be restored by sand-papering. The parts so treated show spotted; even a staining-coat will not cover them uniformly. Fine woods are injured the worst.

CHAPTER XXIX.

FERRIC-PAINT TESTS.

Objection is made by some engineers and paint manufacturers to the immersion methods of testing paints; that they do not meet the actual conditions of coatings exposed to weather; that a ferrice structure is not always wet, but wet and dry, with more dry hours than wet, etc. This would depend altogether upon the location of the structure; in many instances there might be more wet or damp hours than dry ones. A fog or long-continued sweat is more destructive to a paint coating than a passing storm. But the plain fact remains that these tests are all competitive as between different commercial paints, and under uniform conditions. The trial given one paint is given to all; the few successful ones are the better ones to select from to base any subsequent improvements or experiments upon, or for use. The water-test settles the merit of a protective conting in short order, and so soon as generally adopted by those ordering paints for the protection of ferric structures exposed to weather, so soon will the great majority of these patent paint compounds cease to yex the engineer with high claims and low performance.

The nearer any protective coating approximates an enamel or varnish, generally the more durable it will be. The Japanese and Chinese lacquers are varnishes, and dry better by the application of water than in dry air alone, and all compounded varnishes are hardened in the last stages of their drying by water. Lacquers when thoroughly dry remain unchanged for scores of years, when exposed to fresh or salt water either hot or cold or alternately wet and dry, or immersed for years. The coming ferric protective coating will probably be a true

In a paper * read before the Newcastle, England, section of the Society of Chemical Industry, Mr. Henry Smith, F.I.C., described a series of experiments upon the protective powers of twenty seven different English commercial paints, as applied to ironwork in fifty separate instances. The methods of test were those devised and employed by Mr. Max Toltz, C.E., in a series of experiments upon a number of American commercial protective coatings for non-in 1897.1 Three sets of bright and clean iron plates, all of the same size, were respectively coated with the several paints, in all cases furnished as a stiff paste, and when applied, were brought to the consistency of a paint by mixing with genuine boiled linseed oil, capable of drying in seven hours under ordinary conditions of temperature, no draws being used. The first cont was allowed to dry thoroughly firm before the second conting was applied. When this also was firm and hard, one set of the plates was exposed to the weather, as in ordinary cases of painted structures. The other two sets were treated as follows: One set was simply to corroborate the results obtained from the other set, the results being practically identical in each case. Each painted strip was placed in a clean, wide monthed glass battle, half filled with clean pure water. The bottles were not clearl, but were protected from the entrance of dust and impurities while allowing the air free access to the painted plates. Several of the plates, and commenced to corrode in about a week. This was indicated by a cloudiness in the water, which afterward became further oxidized, and formed a red precipitate of ferric oxide, which subsided partly to the bottom of the vessel. After three months' exposure the plates were removed, and the liquid in each bottle, together with the sestiment, was tested for the percentage of iron present in the form of 1956

The figure given as denoting the amount of corrosion is less than the actual amount, as it does not include the partien that adhered to the plate, and was not scraped or brushed off, and would not drain off. In each case the weight of rust was calculated to pounds of rust per 1500 square yards of painted surface; the other figures give the percentage composition of the several paints by weight.

CONDITION AFTER THREE MONTHS' EXPOSURE.

Pounds of Rust from 1500 Square Yards	of S	urfe	ice.		
			C	orros	
			10.00%	Nor	ie.
Red lead	"		12.00%	44	
" "	"	"	7.87%	"	
o sunjuce same owing and of the second	"	«¢	10.00%	Tra	co.
Zine oxide. $.27.27\%$; " $.63.63\%$; " 90.90% ; b	oiled	oil,	9.10%	"	
White lead, pure 92.56%	"	"	7.44%	75	lbs.
White lead, .53.78; barytes 40.33%; "04.11%;	"	"	5.80%	80	"
" " $50.52e_{0}^{2}$; " $42.10e_{0}^{2}$; " $1.02.62e_{0}^{2}$;	"	"	7.38%	95	"
Iron oxide, pale (50%, Fe ₂ O ₂), .83.00%;	"	"	16.40%	81	"
" deep (96% Fe ₃ O ₃), .86.89%;	rnw	oil.	13.11%		"
the the same of the same and th	u	"			
Barytes	• • • • • • • • • • • • • • • • • • • •	• •	11.88%	123	"
Iron oxide, medium volor (94%					
$\operatorname{Fe}_{g}(O_{g})$	"	"	13.11%	124	"
Iron oxide, extra bright color			10.1170	117-2	
$(90\% \text{ Fe}_2O_3),\dots,82.35\%$	"	"	17.65%	127	"
Iron oxide, pure (90% Fe ₂ O ₃) , 76,30%;	11	"	23.70%		
			_		
Barytes	"	"	11.48%	244	"
Indian red $(70^{\circ}_{B} \text{ Fe}_{i}O_{i}), \dots, 82.35^{\circ}_{B}; \dots$	"	**	17.05%	007	"
Turkey red (95°; Fe ₂ O ₃), 81.16°;	**	"	18.84%		
for the state of t			10.04.70	202	
Barytes and calcium carbonate, 62.52%	"	"	10.55%	398	a
Barytes (matural barium sul-					
plate)	"	"	10 0007		- 11
The control of the second terms of the second			12.00%	100	
	c c	"	12.73%	118	"
Burytes and calcium carbonate, 78,80%			•		
from oxide 13.93°	"		422.1.1014		
Harytes and calcium carbonate.60.00°; (••	••	12.14%	242	•
Rose pink (principally barytes) . 12.14%					
Barytes and calcium carbonate 80.50%	u	ti	7.61%	266	"
Colombia pitto			70		
Barytes and calcina carbonate.68.99°					
Ivory and carbon black 8.42% $\left\{ -\text{``}79.87\% \right\}$	boile	d oil	, 20.13%	,352	, 11
Manganese dioxide. 2.46° 1					

Bary(ex and calcium carbonate. 79.30°/ x

mixed with celestial blue, Prassian blue, chrome-yellow, raw sienna, Vandyke brown, Italian ochre, Brunswick and other greens, chromate of lead, English umber, Turkey umber, ultramarine, Chinese blue, burnt sienna, mixed with raw oil in proportions from 11 per cent to 51 per cent of the weight of the paint; the corresion in the order named above ran from 168 pounds to 441 pounds per 1500 square yards of surface.

Except in the case of the blues, umbers, siennas, etc., where the pigment had but little influence on the oil to resist decay beyond that inherent in the oil alone, the more separate substances that entered into the composition of the pigment, the more unreliable it became. A single exception is noted in the case of a Venetian red paint, unde from barytes, calcium carbonate, and a small amount of iron oxide, that gave a better result than barytes alone, or when barytes was mixed with the other color pigments of much less specific gravity. Several substances in a composite point are generally fatal to its protective qualities, no matter to what it is applied. The several atoms of these substances, even if uniformly distributed in the pigment in the process of grinding, bolting, and a ixmy that they are not), will not retain their juxtaposition when mixed with the oil, The heavy atoms will sink, and there will be a marked difference in the coating spread from the top of the paint in the pot from that in the middle or bottom; the lightest and most perishable substances will get on the surface firs Barytes worked well with red lead and zine exide, there being but

Barytes worked well with red lead and zine exide, there being but a small difference in their specific gravities as compared with barytes and the other color or base pigments. With white lead, as the percentage of barytes was increased, so was the corresion. Aside from the reduction in cost of these lead and zine pigments by the addition of barytes, there is no reason for its use, as the barytes alone did not give a satisfactory test. No doubt from the splintery character of its atoms, as has been before commented upon, it is wholly destitute of covering or coloring power. The vagaties of the iron exide paints in the varying proportions of the pigment and oil are noticeable, but not so marked as where barytes, one of the heaviest of all pigments, and calcium carbonate, one of the lightest, both classed as inert

superior to raw oil as a vehicle for the several iron-oxide paints in the ratio of one to nearly five.

Smith's Dish-tests of Paints.

A second series of experiments were made by the same experimenter, and following the method of Mr. Max Toltz, C.E., to wit: A number of iron dishes five inches in diameter and one-half inch deep were scoured bright, and then coated with two coats of the several paints used upon the above-detailed iron plates and under the same conditions as to the composition and drying of the paints. These shallow dishes were filled with water and allowed to completely evaporate in the open air of the laboratory. This operation was repeated six times in the course of six months. Thus tested, the only paints which remained practically unaffected were red-lead and orange-lead paints, some of which, however, such as the "vermilionette" and searlet-red paints, contained a'so a proportion of aniline colors, while two of the red-lead paints contained in the one case 45 per cent of barytes and in the other 66 per cent. All the other dishes were more or less rusted, the order of merit of the better paints being as follows:

1st. Zine oxide.

2d. Equal parts zine white and barytes.

3d. Zine white, 3 parts; barytes, 7 parts.

4th. Lithopone (a mixture of zine sulphate, zine oxide, and barytes).

5th. Pure white lead.

6th. White lead, 5.37 parts; barytes, 4.03 parts.

7th. White lead, 5.05 parts; barytes, 4.21 parts.

All the other paints, thirty-six in number, proved inefficient. The first to show rust was that one painted simply with linseed-oil. The above classification of merit is by Mr. Smith, and, taken together with the detailed report of the glass-bottle test (before given), may be considered a fair representation of the protective qualities of the hundreds of commercial ferric paints foisted upon the market under various trade-mark names in the United States as well as in England,

or even a wooden one. The dish-test, probably, is the nearest to the actual condition which a paint must withstand. When the water in the dish is nearly evaporated, there remains in the circular seam of the bottom a film of water which contains the carbonic acid and the decomposing gases and dirt from the atmosphere, which act upon the paint in such a way that the conting at that part is soon permeated and rust forms. This action is more and more developed after each evaporation, and practically covers the whole dish in a short time. In actual service the same thing will happen. The corner of the dish finds its counterpart in every corner of a ferric structure where two plates, angles, or other parts join. Rust will commence at those seams and extend under the paint, but will not show as plainly on a bridge-truss as on the small dish.

Toltz's Tests of American Commercial Ferric Paints.

The shallow-dish tests by Mr. Max Toltz, C.E. (before referred to), were made prior and during 1897, and extended over a period of from six months to two years. Without entering into as great detail as that quoted from Professor Smith, the deductions from his tests are in brief. Twenty-two different paints were submitted to test under the following classification:

- No. 1. True asplintly variesh paints compounded by heat in the same manner as a black baked japan, and practically of the same nature and comparable therewith. No corrosion reported after the dishes had been filled and evaporated naturally fourteen times.
- No. 2. So-called asphaltic varnishes, or paints of inferior qualities to the above No. 1, made from asphaltum dissolved in benzine or other volatile vehicle, but were not a true varnish. They contained about 43.5 per cent of vehicle and 56.5 per cent reported to be asphaltum. As a rule they showed well in the beginning, but after the volatiles had evaporated, especially when subjected to a moderate heat-test, the coatings became quite brittle, were easily removed by abrasion, and did not protect the surface covered with them. Their composition varied in the several specimens tested. One sample

which they are closely related. The dish painted with only one coat showed a little deterioration at the end of the fourteenth evaporation, while the dishes painted two coats were uninjured, the coating being as clastic and tough as when first applied.

No. 4. Iron-oxide paints consisting of more or less iron oxide with more or less silicious matter, and compounds of lime and magnesia. They were of different grades and qualities, were as a rule well ground and spread well. Under test on the dishes painted with one coat, after the fifth exposure many rust-spots appeared. Those painted two coats were refilled six times, and on them the rust was plainly discernible to the eye.

No. 5. Graphite paints and silica-graphite compounds. These paints were received from the several manufacturers in the form of a stiff paste, and when mixed, ready to apply, 4½ parts of paste to 3½ parts, by weight, of boiled linseed-oil were used. The dishes painted with one coat were evaporated ten times. After the fifth evaporation a few specks of rust were noticeable, and the number gradually increased after each successive evaporation. After the tenth exposure some slight difference between them was noticeable, but not much. The dishes painted two coats were exposed thirteen times in two years, and none of them showed any rust or indication of rust. The natural toughness and elasticity of the paint still remained.

It will be noted that there is a wide discrepancy in the results of the dish-test of Mr. Toltz, as above, of the graphite paints, both the natural amorphous pigments and the compounded silica-graphite pigments, and the plate-test given by Professor Smith of pure flake-graphite mixed with raw linseed-oil that gave 215 pounds of corrosion to 1500 square yards. This, no doubt, is due to the repellent nature of the pure flake-graphite; the pigment does not take kindly to the oil, any more than soapstone does. Raw oil, even if pure, contains from 5 to 7 per cent of water, that renders a combination of the graphite and oil quite uncertain unless under the influence of heat. The boiled-oil vehicle with pure flake-graphite, used by Professor Spennrath in his experiments (hereafter referred to) with paint-skins detached from the matal surfaces, withstood an exposure in a pure water-bath

in daily practice elsewhere.

United States Navy-yard Paint Tests.

The result of these tests corroborate the series of tests made by order of the Secretary of the United States Navy in 1884-5.* request, sixty paint firms submitted seventy-five different paints for test, which were applied to five hundred test-plates, and then immersed in sea-water at four navy-yards, and upon one government. vessel in service. The paints that successfully withstood the test and received an order of merit, were red lead, zinc oxide, carbon, and graphite compounds. The so-called asphaltum paints were at the bottom of the list in the no-merit column. Evidently there has been slight improvement, if any, in this class of paints since the date of the U.S. Navy tests to the present time, and one can but wonder, in the face of repeated and recorded failures, that they ever receive an application to a ferric structure, ashore or afloat. Lead, zinc, carbon, and graphite compounds maintain their supremacy for government work. In other tests of commercial and special paints, where the tests have been carried to the destruction of the coating as a whole, the partial destruction of the vehicle was generally followed by the disintegration of the weaker substances comprising the pigment, such as the carbonate and sulphate of lime, asphaltum, iron oxide, and the various color pigments, viz., the ochres, umbers, blues, greens, carmines, yellows, etc. The only pigments practically unaffected by the destructive element were the graphites, the silica, barytes, slag, slate, and brickdust. Other adulterants were but little affected, some of them being partly recoverable, which was also the case with the red lead, white lead, and zinc-oxide pigments.

Commercial Coal-tar Paints.

Fourteen commercial paints, principally of the coal-tar and asphalt class, were tested under uniform conditions, viz: Wrought-iron plates, free from mill-scale, were coated with two coats each of the following paints. All of the coatings were perfectly dry before the second coat was applied. When the second coats were dry and

bustion gases from locomotives and coke ovens reached them freely at all times, this exposure being similar in all respects to that of rail-way-bridge paints.*

No. 1. Carbonizing coating (Cohen Mfg. Co.).

Physical properties.—Very black, good body, spread well, and covered a large surface, coating smooth.

Drying properties.—Poor.

After 24 hours, wet.

'48 '49 quite wet.

'72 '49 slightly wet.

'49 dry.

Physical test.—In sea-water, much rusted and blistered, paint easily rubbed off, and in bad condition. Atmospheric exposure, condition of coating, fair.

No. 2. Durable metal coating (Edward Smith & Co.).

Physical properties.—Brownish color, thin and required a large quantity to cover; adhered poorly; coating thin and uneven.

Composition.—Linseed-oil, asphaltum, and kauri-gum Drying qualities.—Dried slowly.

After 24 hours, wet.

" 48 " tachy.

" 72 " slightly tachy. " 96 " dry.

Second coat dry in 7 days.

Physical tests.—In sea-water, much rusted, peeled off easily, and blistered.

Atmospheric exposure.—Rusted badly on edges of the plate.

No. 3. Turpentine asphaltum (C. A. Reeves & Co.).

Physical properties.—Coating thick and uneven; required a large amount to cover. Spread poorly; adhered well.

Drying properties.

After 24 hours, quite wet.

48 '' tachy.

72 '' almost dry.

484 '' dry.

spots; did not rub off as easily as No. 2. In two spots corros had *caten through* the plate.

Atmospheric exposure. - Condition very bad, rusted all over.

Physical test.—In sen-water, much rusted, paint peeled off

No. 4. "B." Black varnish (Mica Roofing Co.).

Composition. A coul-tur product, contained light oils, naplatene and authracene, not indicating much pitch.

Physical properties. Very fluid, gave a smooth coating.

Drying properties.—

After 24 hours, wet.

" 48 " quite wet.

'' 72 '' slightly wet,
'' 96 '' dev.

Physical test. In sea-water, fairly free from rust, but cont was rough and uneven, broken off in many places.

Atmospheric exposure. Rusted badly along the edges.

No. 5. Asphaltum paint (C. E. Mills & Co.).

Composition. Asphaltum, petroleum, and some linseed-oil.

Physical properties. Thickens on exposure to air; gives a thuneven cont.

Drying properties.—Dry in 56 hours.

Physical test. In sen-water, much rusted and blistered, la part of the plate coating entirely gone.

Atmospheric exposure. -Plate in very bad condition.

After 24 hours, slightly tachy.

'' 36 '' almost dry.

'' 48 '' '' dry.

Physical test.—In sea-water, did not blister, rusted considerably, paint off in spots. Not so good condition as No. 4, but better than Nos. 1, 2, 3, and 5.

No. 7. "A." Varnish (Mica Roofing Co.).

Composition. Asphaltum, in petroleum spirits.

Physical properties.—Fluid, spreads well and adheres well; gave a thick coating fairly smooth, smells strongly of petroleum spirits.

Drying properties, |--

After 24 hours, quite wet.

'48 '48 slightly tachy.

'460 '4 dry.

Physical test.—In sea-water, much rusted, not blistered, paint off in numerous spots. In bad condition.

Atmospheric exposure. - Badly rusted.

No. 8. Mineral rubber (Assyrian Asphalt Co.).

Physical properties. This paint was so thick and viscous that it could not be applied without thinning. When thinned with naphtha it did not work satisfactorily, and the experiments with it were abandoned. (See Chapter X11.)

No. 9. Black roofing paint (Samuel Cabot).

Composition. Pitch dissolved in light petroleum oil.

Physical properties. Fluid, gave a smooth coating that adhered well. Smelt of tar-oil.

the time of the control of the contr

Drying properties,—Dried in 56 hours.

No. 10. Black paint (Thomas Mfg. Co.).

Composition. A coal-tar paint with a heavy oil menstraum.

Physical properties. Much like No. 9.

Drying properties.

After 24 hours, wet.

" 48 " tuchy.

" 60 " dry.

Physical tests.—In sea-water, paint came off easily; many rust-spots.

Atmospheric exposure, -Condition fairly good.

No. 11. Slay cement paint (Burrett Mfg. Co.).

Physical properties. A coal-tar paint, producing a coating similar to Nos. 9 and 10.

Drying properties. Dry in 60 hours.

Physical tests. In sea-water, had a tendency to peel off; some rust-spots noticed.

Atmospheric exposure. Plate bully rusted.

No. 12. "Ferrodor" (Wm. Somerville's Sons).

Composition. Graphite, turpentine, exide of iron, and linseed-oil. A compound or patent paint.

Physical properties. Color, purplish gray; conting, very thin; three conts recommended by the manufacturers.

Drying properties. Dry after 48 hours.

Physical tests. In sen-water, paint peeled off badly, plates very much corroded, bad condition generally. The graphite settled to the bottom of the can in a tenacious pasty mass, and the paint was spread with great difficulty.

No. 13. "Antoxide." Ready mixed paint (Harrison Bros.).

Physical manuating - Briefit rul miles dece to mil tout

Composition. A paint made from coal-tar (special process). Physical properties.—Deep-black color; spread well and adhered

well, giving a smooth even coating, rather thick; contained 10 per cent free carbon.

Drying properties. -Dry in 36 hours.

Phusical tests. In sea-water no corrosion or blistering; had a slight tendency to peel.

Atmospheric exposure. -Plate slightly rusted; stood the action of combustion gases better than any of the other competitive paints. There was no tendency of the paint to run or crawl when applied to any metallic surface at ordinary temperature. (See Water-pipe Contings, Chapter XII.)

A number of commercial ferric paints were tested (1899) by Prof. Im O. Buker for their comparative resistance to heat, seawater, strain, elasticity, the fumes of sulphuric and nitric acids also carbonic acid, with the following results:*

The samples of paint named were furnished by the manufacturers for the purpose and mixed with linseed-oil as directed by them, and sprend on elean bright wrought-iron test-plates.

Reference Number.	Kind of Paint.	Weight per Callon.	How Received from the Maker.
1 2 3 4 5 6 7 8 9 10 11	Red lend White lend. Purple iron oxide. Chattanooga iron oxide. Williamsport iron oxide. Detroit superior graphite, Mexican graphite. Dixon's graphite. Trinidad asphalt. Bessemer point. Carbonizing conting Lithogen siliente.	Pounds. 31.72 21.24 11.34 14.13 12.00 9.40 8.57 8.84 9.52 13.00 9.61 17.06	Dry pigment, Paste, Mixed ready for use, Dry pigment, Mixed ready for use, """ Dry pigment, Mixed ready for use, """ """ """ """ """ """ """ "

A sample of each paint in one and two-coat work was exposed to heat and the products of combustion in the smoke-flue from a boiler burning bitumnous coal. The condition of the plates on removal was:

The powdery resi-1 Red lead - Entirely dead and very brittle.

entirely dead, and very easily removed. Itemoval of the residue exposed the base metal.

3. Purple iron oxide. Covered with blisters. Paint soft and

easily removed. Removal of the blisters exposed the base metal only to a slight extent.

4. Chattanooga iron oxide. Considerably blistered, moderately soft, adhering well. Removing the blisters did not expose the base metal.

5. Williamsport iron oxide. Covered with blisters. Paint tenacious, adhering well. Removing the blisters exposed the base metal.

6. Superior graphite. Small blisters. Paint hard and difficult to remove. Almost impossible to expose the base metal.

7. Mexican graphite. Soft and readily removed. Removal of blisters did not expose the base metal.

8. Dixon's graphite. Very few blisters. Paint hard and very adherent.

9. Trinidad aspludt. Smooth; considerable of the paint melted and run off. The portion remaining was hard and brittle.

10. Bessemer paint. Soft and hard to scratch off.

11. Carbonizing conting. Ridges very corepieuous. Paint firm and adherent. Removal of blisters exhibited a very porous, loose structure of the paint.

12. Lithogen silicate (white paint). Substantially the same as the white lead.

Effect of salt water. A set of the plates were exposed to a saturated solution of sensalt (brine) for seven weeks, the plates being frequently withdrawn and allowed to dry. Their condition at the end of the exposure was:

1. Red lend. Hard and adhering. Metal clean and bright under the two coats, but badly rusted under the single coat

2. White lead. Hard and adhering. Clean metal under two coats, rust under the single coat.

3. Purple iron oxide. Firm and adhering. Occasional rust spots throughout.

4. Chattanooga iron oxide. Firm and adherent. No rust anywhere,

under the single cont, none under two coats. 7. Mexican graphite. Elastic and easily removed. No rust.

8. Dixon's graphite. Good condition. Elasticity only slightly impaired. Easily removed no rust.

9. Trinidad asphalt. Seemingly unaffected. No rust anywhere. 10. Bessemer paint. Moderately hard, peeled off easily, no rust.

11. Carbonizing coating. Peeled off easily. Rust beneath both

the one and two coats. 12. Lithogen silicate. Hard and adherent. No rust-spots.

No report was made of the condition of the paints exposed to atmospheric influences, evidently for the reason that the exposure period had not been long enough to materially affect any of the paints when the test closed.

A set of the plates were exposed to the fumes of strong sulphuric-, nitrie-, and carbonic-acid gases for five weeks. The action of the sulplanrous gas was characterized by its bleaching power upon the paints and the disintegrating effect on the iron under the paint, also the formation over the entire area of blisters, under which was found a moderately hard whitish deposit.

The order of merit for the several paints was: Trinidad asphaltum, Carbonizing conting, Dixon's graphite, Red lead, Lithogen silicate, Mexican graphite, Purple iron oxide, Superior graphite, Bessemer

paint, Chattanooga iron oxide, Williamsport iron oxide, White lead. The effect of the nitrie-acid gas was substantially the same as the sulphurous gas, except that the single contings of the paints were completely destroyed. The order of merit for the double coatings was: Trinidad asphaltum, Lithogen silicate, Red lead, White lead, Purple iron oxide, Superior graphite, Mexican graphite, Chattanooga iron oxide, Williamsport iron oxide, Dixon graphite, Carbonizing coating, Bessemer paint.

The earbonic-acid gas in large quantities, supplemented by moisture, had only an almost imperceptible effect upon any of the paints.

To test whether any of the paints would crack during the clongation of a painted bar, strips of machine steel $2'' \times \frac{1}{2}'' \times 18''$ were painted two coats, and after drying for two months were submitted to a strain

of 16,000 pounds per square inch; the paint in every case remained e t t t tt t the two to the their increased clearly It was noticed that after passing the elastic limit of the stathe paints were marked by a series of lines arranged in herriabone patterns, that were alike on both sides of the bar and al situated. Evidently the lines were due to a tearrangement of atoms of the steel bars while under strain, the centre atoms moveless freely than those near the corners and edges of the bars, a paint naturally following the particles of steel that they cover Naturally the heavier contings of the paints were the least class. The order of merit in the elasticity test was. Trunched a phaltu Carbonizing conting, Purple iron oxide, Dixon graphite, Mexicographite, Superior graphite, Williamsport iron oxide, Chattanon iron oxide, Bessemer paint, White lead, Red lead, Lithogen sheater

CHAPTER XXX.

PAINT TESTS ON RAILWAYS.

New York Elevated Railways.

The physical condition and the extent of corrosion on all parts of these structures have been freely commented upon by the technical engineering journals and the daily press. When originally erected, no attempt was made to remove the mill-scale, and none has been made since, presumably because of the impossibility of its success, and the cost. The composition of the paint which has been repeatedly applied to them has been kept very uniformly good in quality, but its application has been solely for appearance, as no paint can now reach the seat of corrosion underlying all the coats of scale, dust, cinders, and paint.

The renewal of the whole structure will probably be necessary in less than a hundred years from its erection.

The composition of the paint used is given by the chief engineer as follows:

FOR FIFTY GALLONS OF PAINT, OLIVE-DRAB COLOR.

				Summer Formula.			iter iula.
oest				300 r	oounds	275	pounds
, strictly	best o	_l uali	ty	175	"	150	"
"	"	"		100	"	90	"
"	"	"		1	"	1	"
"	"	"		3	: "	•	<u> </u>
	, strictly	, strictly best o	, strictly best quali	s, strictly best quality	best	best	best

5764 pounds 5164 pounds

The above pigments are ground in Campbell & Thayer's raw linseed-oil. The weights given include the necessary oil to grind the pigments to a paste.

when applied, it was mixed with				
Boiled linseed-oil, Campbell & Thayer's	8	gall.	9	gall.
Raw linseed-oil, " " "	15	"	15	"
Spirits of turpentine, first quality	3	"	3	"
Liquid or japan drier, " "	2	"	3	"

New York Elevated Radway Vandact

The viaduet over the Harlem Station of the New York Elevated Railway at 155th Street was oil coated, and received iron-oxide paint coatings at the time of its erection, and within the years of its completion had developed corrosion to such an extent that in 1897 the sand-blast was used to clean it preparators for another effort for its preservation. This sand blast process cost about \$10,000, or over lifteen cents per square foot to apply, or about seven times more than a properly selected method of procedure and paint would have cost in the first place, and then only the lower and accessible sides or parts in sight received treatment. About 50,000 square feet of surface was cleaned by the sand-blast, to the bught iron, removing about 12 tons of old paint, scales of rust, and conders, showing a unimber of distinct layers of highly corroded matter.

Seventeen panels of lattice-truss, these beam and buckle-plates. supporting the paved carriage readway and feetpaths excrheal. about 2825 square feet of surface each, and numbered consecutively I to 17, were then painted with the same mumber of selected protective contings furnished by a like must be of peaks firms in competition with each other. The neveral coatings were applied in strict conformity to the directions received with each brainly of passet, the appliention being to the bright from an left by the action of the nated blast, and within I to 4 hours from the time the based blant commed action, Every tuned like condition was broadingle sale becausing the sounds the test one of a practical and commercial mature as well as est seneratific value, absolutely without prejudice or favor in any respect. From the premification of the otherwise as as arragarance and and and and the others into expended to starting sea mir, for, cambers, steam, and games from scores of locomotives in constant service beheath it meanly all of the metal being within a few feet of the tops of engine stacks and receiving the produced the contibustions are force to ever to the first error and the thirty of the product of Classed state of the follower transles as an expansion house, and five an expansioner

perfect com	lition of t	he coating.
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Rate of Paint. Rate of Drying.	Freedom from Rust, Per Cent.	Reference Marks as to Condition.
1 3 Lead, graphite and Incol-oil	97 80 25 75 99 85 75 30 80 78 58 92 67 70 65	a b c d e d d f a g h a d j k l

- a. Very little rust. Paint crumbles in places as though rotten. Easily removed.
 - b. Fair condition, but discolored; rust coming through,
 - c. Very badly rusted,
 - d. Rusty, but not deep,
 - c. Slight rust on top flauge of one girder; rest of girder clean,
 - I. Rust very deep; buckle plates had,
 - g. Aren of rust spots small; rust not very deep,
 - h. Rust very bad and deep.

 - j. Deeply rusted; buckle plates still good.
 - k. Rust very deep und angry; buckle plates mildewed.
 - 1. Small pumples of rust, as though formed under the paint.

Panel No. I was an outside one, and the first to be sand-blasted and painted, in some parts with two and in others three coats of paint, in the clear hot days of summer, a material advantage in its favor. The sand-blast was then shifted to the southern end of the vinduct; and panel No. 17, also an outside our, was the next one cleaned and painted in hot clear weather, and so on consecutively, in

the reverse order of the panel numbers, back to No. 1; panels Nos. 7 to 2 having been done late in the fall under unfavorable conditions as to the spreading and drying of the paint in addition to the other objectionable conditions. About 80 square feet of panel surface was cleaned per hour, or 600 square feet per working day; each panel requiring from five to six days to clean and paint it.

At the end of about a year the condition of all of the paints was so unsatisfactory that the vinduct was repainted without removing, only in a perfunctory manner, the old test coatings with their fast-forming hurdens of rust; and this competitive test came to an inglorious end,

The result could have been foreseen from the first, before a single truss or pound of material had been placed in position, or was even out of the construction shops, bad not conducted great, official indifference or ignorance, either one or all, ruled the matter

The destruction of the tubular railway bridge over the St. Lawrence River at Montreal, Canada, had not become a fact so masty with age as to have escaped attention concerning the dangerous effect of hot combustion gases upon any paint continuous as a confined space, Corrosion history blindly repeated itself when the viaduet material was first pointed by the contractors, then repeated the "Connedy of Errors" when it was creeted and again when it was sand bla red for its final flasco. The plain facts of the painting after the sand blast action are, that the contags were destined for an early destruction from the beginning, by reason that the first coat was applied in an atmosphere saturated with the hot vapors of combination and steam, which were so corrosive that the freshly cleaned surface of the metal showed a blash of rust within an hour after cleaning, and if left for three hours the rust could be wiped off by the hand. The paints were spread in this atmosphere, and before they could in any measure dry, so as to be in any degree resisting, they were thors oughly impregnated by the hot gases and steam which left their condensed strength upon the surfaces of the green paints. The second and subsequent conts were not only applied under the same atmosA paint test of an extended character has been in progress for the past few years by Mr. Geo. W. Welster, C.E.,* to determine the best paints for use on the city street iron bridges, crossing the railroads within the city limits of Philadelphia, Pa.

Fifty-four sample plates of iron 12"×24" were coated by twenty-two manufactures of paint, and exposed at a number of places on the street viadacts, in situations that were as nearly uniform for the several competitive coatings as possible to provide. The test coatings were changed in their location as circumstances required to equalize the exposures, which were very severe, the clearance between the top of the locomotive stacks and the metal work of the bridges being only two to three feet.

The samples of paint submitted included the most prominent proprietary paints, including the earbon and graphite classes.

The results of only a few months' test demonstrated that on the lower surfaces of the plates on the bridge structure no paint was able to resist the mechanical injury from the sand-blast action of the locomotive exhaust. These situations are now protected from this action by wooden sheeting a few feet in width on the line of the exhaust.

On the upper side of the test-plates, subject to moisture, combustion gases, and deposits of einder, the results were more satisfactory, but were not conclusive as to the relative merits of the samples, due to the difficulty in comparison on the basis of truly identical conditions.

The general trend of the results was, the subsequent selection of certain of the proprietary paints for a trial on bridges, and component parts of bridges, under a general formula, having red lead as the principal pigment, viz:

Red lead, two coats over shop coat of raw linseed-oil for inclosed space of structures.

Red lead over shop cont of raw linseed-oil and two coats of white lead three parts, and zine oxide one part, for the field work.

Indian red, one cost over shop cost of oil. Red lead, third cost.

^{*}Chief Engineer's Bureau of Surveys, Department of Public Works, City of

and the second s

Indian red, one cont over shop cont of oil. White lead three parts, zine oxide one part, for field work.

The Gray's Ferry deck bridge over the Schuvlkill River was included in the test, being painted with the following paints, all applied as a first or shop coat:

Nobrae, lucol-oil paint, rubber paint, Bessemer paint, antoxide, durable metal coating, red lead. Above the deck for the second full coat, red lead was applied, except in one case, where white lead and zine oxide were used. Below deck, for the third full coat, the same paint was used as for the shop cont, except in one case where white lead and zine oxide were used for both the second and third coats.

The above contings were applied in 1899-1990, and the time since then has been too short to note any material difference in their condition. The general practice in painting ferric structures by the Board of Public Works of the city of Pluladelphia for a minder of years has been the use of red lead and lamplified for a first or prime ing coat at the shop, followed by two coats of the same paint in different shades of chocolate color for the field coats, though in some cases white lead and zine oxide have been the held coats. That the above paints have not proven satisfactory in the presence of combustion gases and other influences incident to their location is exident from the above experiments to correct their deterencies. In connection with the same matter, it may be of interest to note that the train-shed roof of the Broad Street Station of the Pennsylvania Railroad at Philadelphia, which was painted with red lead and lamp black, is seriously affected by the corrosion of the root trusses. This structure has had extremely good care since its erection, but corresion has established itself, owing to the early decay of the red lead coatings, and will soon require eleming by the sand blast to correct the mistake of using red lead for train-shed painting time Chapter XXXVI, Changes in Pigments.)

Influences that Affect Paints

Some of the influences that affect the life of a paint conting have been determined by the experiments of Prof. J. Suggestable from

dilute solution of sulphuric acid and the zine dissolved. The paintskins were then used for testing by immersion and exposure for six months in a number of liquids and gases, as follows:

Immersion Tests

In pure rain-water the skin remained cohesive, even clastic, was of dull color, noticeably injured, and			
In sea water the skin remained uninjured in tex- ture and lustre, with a small loss in elasticity, and in	1().4	per	cent.
h a 10 per cent solution of common salt the	4.52	. "	"
skin was but little affected in histre and elasticity, but lost in weight	2.4	u	"
skin was unchanged. Lost in weight	3.5	tt	"

1.65

9.83

"

In an aqueous solution (alkaline) of mineral-coal ashes the skin was materially affected. Lost in

In a 5 per cent solution of nitric acid the skin was destroyed, In a 10 per cent solution of the chloride of mag-

In a 10 per cent solution of the chloride of magnesium the skin was unchanged, but lost in weight

Exposure Tests in Closed Vessels,

Over sea-water for six months the skin was uninjured in color or texture, but had become somewhat viscous: no loss in weight.

Over acctic acid, funning nurriatic acid, nitric acid, ammoniacal liquor, liquid sulphate of ammonium, a solution of gaseous sulphurous acid and water, all the skins were destroyed in a few days.

A skin made from red lead and linseed oil, exposed for forty-eight hours to an atmosphere of hydric sulphide, became black and rough, dull in lustre, and increased in weight.....

15 per cent.

The changes here indicated relate solely to the vehicle, as the graphite pigment was passive to the action of any of the destructive agents. Wherever the skins were destroyed every other oil-paint conting would have been likewise destroyed, whatever pigment was in it. In the other cases where changes in the vehicle are noted, the change of the skin appears to be wholly unlike that which would have occurred had it been attached to any surface. The professor has evidently found this to be the case, judging from some of his notes preceding the record of his tests. There are commercial paints notably those made from amorphous mineral graphite, that containing less graphitic carbon, and combined with silica and a small amount of mineral oxides—that would have afforded a better protection to the vehicle than the chemically prepared graphite used in these experiments, the cost of which would probably bur it from forming any part of a protective covering for non-

Spennrath's Temperature Tests.

A number of graphite paint-skins of the same character as those used in the above immersion and exposure tests; also, some three conted skins made with other pigments and linsect oil and mixed with turpentine and other driers, and mineral oil, were submitted to constant temperatures of 122°, 203°, and 248° F, for five days. Briefly the results were:

All the skins shortened from 1.2 to 4.3 per cent, averaging 3.76 per cent, and lost in weight from 2.11 to 8.3 per cent, averaging 5.52 per cent, averaging

was changed the least in other respects. All the other skins became brittle and stiff, broke easily when bent sharply, and were darkened in color. The white-lead skin changed to a faint yellow, the sulphide of lead and zinc oxide skin to an intense yellow.

The addition to the linseed oil of 10 per cent of either the oil of turpentine or other driers, or a mineral oil, or other fatty non-drying oils, including some gum copal, had no effect whatever to resist the changes effected by the heat. Not more than 10 per cent of mineral oil could be added to the linseed oil, as it rendered the paint viscous after drying. The Bessemer paint-skin, the pigment being a ground furnace slag, and the linseed-oil vehicle having an addition of a non-drying fatty oil with some gum copal, was just as sensitive to the heat as any oil paint.

Generally the action of the heat was less marked upon the graphite skins, which were less brittle than those made from white lead or zinc-white. The red-lead skins were especially sensitive to mechanical influences.

These changes are easily accounted for. The oily, repellent nature of the flake graphite prevented it from bonding to the oil vehicle as firmly as the other natural pigments and those of higher specific gravity will do in both a green or a thoroughly dried paint. While it is generally known that heat is destructive to oil-paint coatings, it does not follow that the coatings are so sensitive to its action that it may be deemed the principal cause of their failure. The engine and fire-rooms of ocean steamers and war-vessels are exposed to temperatures of 120° to 140° F. for months at a time, and many times in succession in atmospheres heavily charged with moisture and other vapor, without any material disturbance to the protective character of the coatings.

There are commercial paints in extensive use, subject to temperatures of 300° to 350° F. under pressures of steam, which preserve their integrity after years of exposure. In these instances the life of the coating depends quite as much upon the vehicle as upon the pigment.

CHAPTER NNNL

PAINTING BY SPRAY.

PAINTING by spray or the air-brush has lately come largely into use; in fact, it would have been impossible to have covered, in any acceptable manner, the World's Fair Buildings erected since 1890, without the use of the paint-spray process.

At the Columbian Exposition, the results of the spray method of painting, compared with the use of the hand-brush, were: A corps of hand-brush painters, working in the usual manner of applying kalsomine, averaged about 800 square feet of surface daily, while 16,000 to 20,000 square feet were covered by a spray-machine in eight hours, 30,000 square feet having been reached under favorable conditions. In the Mannfacturers' Building, with a daily average of thirty men using spray-machines, at the end of eighteen working days, 1,332,700 square feet of surface had been covered; equal to an average of 2368 square feet per day per man. This was during the coldest days of winter, when the water paint, in attempting to spread it by hand, froze solid. The spray required about twenty one gallons of kalsomine against twenty gallons by the brush, but the saving in labor was nearly twenty to one in favor of the spray process.

Gas-holders in duty are exceptionally lard to paint. One painted by spray, using iron-oxide paint, averaged from 2700 to 2900 square feet of surface per hour for three men using two sprays. The amount of paint used was not notably more than with the brush. The cost of the labor was less than 4 cent per square yard.

In the Michigan Engineers' Manual for 1897, Mr. J. J. Huber describes an extemporized spray-machine, and the results in painting 100,000 square feet of rough heudock siding with iron-oxide and raw-oil paint.

The contractor's bid for labor, ladders, and brushes, the company to furidsh the paint, was Cost of the paint, labor, and apparatus, lifteen cents per 100 square feet, or less than one-half that of painting by hand. The paint could



Fig. 41.—Field spray apparatus at work.

be applied from eight to ten feet above the spraymens' heads, and ordinary laborers could do the work.

In some experiments made by the P. & L. E. R. R., using a spray apparatus for painting box freight-ears, the time required was thirty minutes per ear; one man with an eight-inch brush following the spray thirty minutes more, or total of one hour per ear for each coat. To paint a 60,000-pound-capacity coal-car required two men twenty minutes each, spraying the lettering not included.

At the Master Car and Locomotive Painters' Association, in 1897,

time and costs in detail of painting box freight-cars by brush and spray.

	With th	n Brunli.	With the Spray,			
Work Coated.	Time.	Cost.	Time.	Crist,		
Sills, one coat	20 min,	\$0.05	13 min,	\$0.031		
Edge board, one coat	40 "	0.10	17 "	0 011		
Body, three conts	7 hrs.	1.05	84 "	0.21		
Puttying up	1 "	0.15	1 hr.	0.15		
Roof, two conts	30 min,	0.073	12 min,	0.03		
Trucks, one cont	1 hr.	0.15	20 "	0.115		
Blacking ironwork	25 min,	0.061	25 11	0.061		
100 A 2						
Totals	rs, 55 miu, -	\$1.637	3 lus, 51 min,	\$11.574		

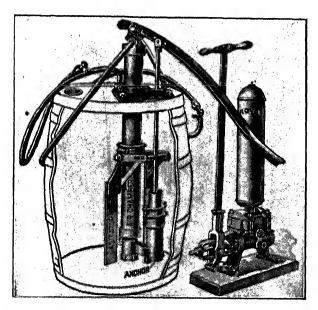
Result 1.98 to one in time, and 2.82 in cost, in favor of the spray. The danger to the health of the painters in the use of the spray is very marked over that in the use of the brush, whether kalsomine, iron oxides, or mixed paints are used. In the spraying of lead paints or those containing any metallic oxides the dangerous effects are greatly increased, even when the greatest possible care is used to guard against them. The fine mist-like spray is readily taken into the lungs at every respiration, and is more thoroughly introduced into the system than is possible by absorption from contact in painting by hand.



Fig. 42.—Mathewson's patent belief for painting by spray or cleaning by the much blant

These were applications from compressed-air installations used for other purposes than painting.

The merit of oil-paint spray coatings has not been fully established. The spray necessarily carries a part of the air with the condensed moisture in it into the paint, and its subsequent escape by expansion and evaporation must result in a more porous coating than with paint applied by a hand-brush. Following the spray immediately with a brush will remove the porosity to some extent. The brushing out of any paint is a great factor in its durability, and as the use of the spray renders the employment of a cheaper grade of lahor more feasible than with the use of the brush, the effects of an indifferent use of painter's "elbow-grease" will soon reveal itself in the decay of the conting.



the paint is most necessary. Peeling of the paint and corrosion promptly follow may extended application of an oil-paint spray coating on a ferric body in situ.

The methods of painting were recently discussed by the Western Association of Railway Superintendents of Bridges and Buildings, in answer to a circular asking for information upon the subject. Eighteen answers were received. All were in favor of the sand-blast for cleaning either new or old metallic surfaces preparatory to painting.

Six were in favor of the air-spray for some classes of work, three were opposed to it, and nine were non-committal. Two who had tried it were opposed to it. One superintendent said: "On iron bridges other than on plate girders I found that there was more paint wasted than applied to the structure. The waste in attempting to paint lattice-truss work was very marked, and the coating was not equally or well spread." He favored the use of a stiff hard brush to insure a close contact of the paint, which he could not get with the spray. Too much nir was incorporated with the paint by the spray, and would not release tself in the drying of the paint. It left the coating more porous than with the use of bandbrushes. Following the spray with a hand-brush did not materially help the coating in durability, when compared with surfaces spread on the same structure at the same time, by the same painters using hand-brushes, and the same paint.

"The use of the spray, following it with a heavy hand brush, was admissible upon some of the large wooden buildings, as the



Fig. 44.—Hand spray apparatus.

failure of the paint in these cases was not attended by corrosion, blistering being the principal cause of failure in the sprayed coating on wooden and masonry surfaces."

Generally, heavy or very thick oil paints cannot be successfully spread by spray, unless under air pressures of sixty or more pounds per square inch. This renders the

use of the spray for oil paint useless, unless power other than hand-

fluidity. If the paint is thinned with benzine or turpentine to the point where a moderate pressure of air will enable the spray to work without choking in the nozzle, all of the objections to this class of paints are increased, as the extra amount of volatiles in them to be evaporated leaves the coating more porous and hastens its decay. Two coats of such air-sprayed paints are required to equal in protecting power one coat of heavy hand-brush work.

With kalsomining or water paints the spray apparatus finds an almost uncontested field, and from the great saving in labor is recommended.

CHAPTER NAXII.

MINIOU PAINTS.

REPUTABLE manufacturers of standard pigments are greatly at the mercy of many of the proprietary or patent paints ready for use that are a feature of the paint market.

Standard pigments subsequently appear in these compound paints mixed with a variety of well-known inferior substances that by some alleged special mechanical manipulation "developed in our factory" makes of them a preminently superior product, "wholly unlike that produced by the antiquated process employed before the advent of our new idea."

Mixed paints for the great bulk of the paint trade are a convenience that cannot be ignored, and are in conformity with the general advancement of the times. Responsible manufacturers and dealers in paints furnish them; and they are more uniform in quality and color, of better composition, as well as cheaper, than when mixed by the individual painter.

Responsible paint manufacturers inspect and test the quality of all their materials, and are certain that they are standard in all respects, more than it is possible for the individual painter to do, however much he may desire to produce a good paint.

Railway companies and bridge-manufacturing firms, from the magnitude of their painted work, are able to employ the necessary staff to secure good materials, also the technical knowledge to mix and apply them. The application of the paint in a great measure is under their control, and the composition of it can be varied to meet all the conditions as they arise, and a direct responsibility established for any failure in the coating.

Failures are possible and occur with the best of paints. The

upon, finding that the paint does not work or spread well, doses it with benzine, turps, or some other volatile or vehicle. He does not always know or care what the composition of either the paint or the added element is, so that it enables him to get through with his work without an immediate appearance of distress in the coating.

Special mixed commercial paints generally require a special order of procedure in their application, and when these are faithfully

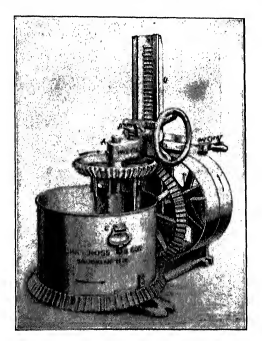


Fig. 45.—Power paint-mixer.

carried out will usually give better results than when the application is left to the ordinary painter's manipulation.

A few special paints that have passed the fortuitous requirements of the United States and foreign patent offices, and have one or more trade-marks to each combination, are the following:

Eira- and water-proof. Composition: coal-tar. oil, gypsum, Japan,

Fire and acid-proof, Composed of coal-tar, pitch, common mineral paint, hydraulic cement, gray ochre, asbestos, slaked line, liquid drier, and litharge.

In most of the above and in other similar patent compounds the quantities of each substance are not defined, but left to the discretion of the user.

Other compounds of a kindred nature contain saltpeter, sulphur, caustic potash, mica, tale, zine slag, salts of tartar, exide of copper, shellae, sulphate of merency and sulphate of zine, verdigtis, copperas, india-rubber, hydraulic-cement slag, sompstone, solutions of gall-nuts, tannin, acetone, yellow soap, lignom vite, garlie, asafetida, and one or more of the list of inert pigments given in Chapter XVIII.

These compounds are recommended as special paints for ferric structures. In most cases the merits are so blindly set forth that one is in doubt whether it is the preservation or destruction of the paint or the covered surface the proprietor wishes to secure,

As mentioned before, all mixed paints are not necessarily objectionable compounds, and to be avoided. A combination of pigments to secure a desired result is often necessary where the use of one pigment would be ineffective.

In the cases of red lead and lampblack the lampblack delays the setting, adds body, prevents the crawl or curdling of straight red-lead coatings, and is in every way beneficial to the physical character of the paint.

There is enough oxidizing element in the red lead to cause the lamphlack (which is a slow drier) to dry without using a large amount of japan or other drier.

A small amount of French order is sometimes added to redlead and lampblack mixtures to give a brighter tone to the charmlate color of the mixture, making what is called the "Pullman color," This mixture has proved to be very reliable under some severe exposures on cars,

The specifications for the painting of one of the largest bridges

in linseed-oil containing a small amount of Japan drier, will outwear red lead. If the coating is applied to a rusty surface, the scales of rust will break through red lead sooner than through the above mixture.

Mixtures of zine-white and white lead, both true pigments, are thought by some engineers to be a more durable coating than either alone, and for some external exposures are said to be improved by the addition of 10 to 15 per cent of silica or barytes.

The burytes and silica in these cases, also when added to flake graphite, save oil and give weight and bulk, as well as a frictional element to hold the graphite in place during the setting of the paint.

In all these instances the paint appears to be better for the presence of the various substances; but could the same group of materials be combined into a single pigment, it would prove superior to any mechanically arranged article.

The few cases where the mixture of true pigments with each other or with an inert substance has proved to be beneficial are not numerous enough to afford any foundation for a mass of incongruous substances called "mixed paints" that flood the market, and which the reputable paint-manufacturer is almost powerless to stem. (See Chapters V. XXX.)

No reliable paint can be made without skilled labor at almost every stage of its manufacture, even with the aid of the best mechanical devices to reduce the labor account. Generally the labor and power account is two and a half to three cents per pound of paint, or twenty-five to thirty cents per gallon. Any paint worth applying to a ferric or any other structure of importance cannot be bought for forty or fifty cents a gallon. Such paint will not protect a ferric body, and if applied will prove (because of frequent renewals) more expensive than one costing three times as much.

Iron oxide is the cheapest straight pigment in use. But this cannot be properly ground in a reliable oil, barreled, and delivered for less than seventy cents per gallon, unless the manufacturer is losing money, or using an adulterated oil or a very unreliable iron-oxide pigment.

to 120° or 130° Fahr, and benzine is slowly added and well stirred. the mixture will not separate on cooling, but remain fixed until the oil is spread and dried by the evaporation of the volatile. The odor of the benzine in the paint in this case is almost suppressed; it is only markedly noticeable when the oil is again heated to near the above degree. Hot mineral oil added to linseed oil is hard to detect by the odor. but the character of the mineral oil as a non-diving oil is not chanced in the slightest degree by the heating. A gill of nametal oil added to a gallon of red-lead paint will delay the setting of the paint. The paint never dries hard, but only on the surface. It temanes viscid beneath, and the conting is liable to preliat any time-There were sold in the United States in the year 1900 (30,000 (00) gallons of mixed paints and pastes, the use of which is metercing about 10 per cent each year. More paint is used in the United

against them. Oil made from unripe, smoky, conden ned, or "nograde" seeds, that often contain almost as much non-drying oil as the drying element, is too frequently used, and benzine is used for the drier instead of japan or turpentine. When line coloul is heated

mints are adulterated quite as much as any painty produced in the United States. The highly extolled English "Torbay" paint is made from a 90 per cent from oxide against, and has no medit over any American brand of oxide paint containing the same percentage of iron oxide

States per capita than in any other country. The Loglish mixed

and an equal quality of American litered oil

"Ferredor," an English trade-mark mixed paint i exploited as being manufactured from a "natural metallic steel may, 95 per cent rustless peroxide of iron found, in a very fine state of division, as a crystalline peraxide, and surpasses the oxides of iron, and is superior to red lead. 'Ferredor' cannot alesale or impart exycu. so that the oil in the paint is not destroyed, as is the ener with the red oxides," etc.

The most brazen advertiser of any grade of American mineral brown paint never equalled this, and no American has been bold enough to attach to his product so fearful a trade-mark as "Schargers by using this firm's product, though it is but a corrosive one.

"Armour-Scale Paint" (Panzerschuppen). A Swiss ferrie paint made from a number of formulæ, also one or more German paints bearing the same trade-mark, are simply iron-oxide paints made from 80 to 90 per cent iron ores that the manufacturers call "granular micaceous." A greasy, crystalline, scaly iron ore with gangue, etc. Graphite in some (nuclassified) form is added and the vehicle is sometimes a linseed-oil varnish.

"Lender's Anti-Corrosive Paint," especially recommended as impervious to heat, cold, warm water, steam, volatile acids, alkalies, gaseous ammonia, hydrochloric acid gas, and sulphuretted hydrogen gas. The base of the pigment is called "a silicate of iron." It is simply an ordinary iron ore containing iron-oxide, 88.56 per cent; silica, 5.40 per cent; lime and magnesia, 3.10 per cent; alum and phosphoric acid, 0.55 per cent; undetermined substances and loss, 4.30 per cent. It is sold in the form of a paste, being finely ground in a holled oil or varnish, and when used is reduced with raw linseed-oil, and litharge added for a drier. A special point claimed for this wonderful oxide-of-iron paint is that "any mineral paint of the right color can be added to produce the desired tone." Wonderful product! The special advantages for this special paint are accompanied by a specially high price for it.

Other examples of the foreign mixed paints and the art of advertising them could be cited. A mixed paint is not necessarily a good one because of its trade-mark and high price, nor is it a notoriously had one because of its being an American firm's product.

Some instances of the unreliable character of compound paints have been given in Chapters V (White lead) and XXX (Paint tests). The following instance, from the cost and magnitude of the structure to which the paint was applied, the public interests at stake, and the result of the application, is of interest. It illustrates the unreliable character of compounded pigments applied over other basic pigment paints for the protection of ferric structures.

The new suspension bridge over the East River has in Brooklyn a viaduct about 9000 feet long. This viaduct is of lattice trusses

of boiled oil before machining. After riveting up in sections preparatory to shipment it received a conting of red lead paint, and after erection another conting of red lead and knopletack paint was applied.

After a number of months' drying the first tair lairy coat of Jewett's white lead, 70 to 75 parts, and the New Jevey viae coate 20 to 25 parts, ground in raw linseed oil, was applied. The engineer corps, from some previous experience with this maxture, were particular to see that the quality of all the material was standard in all tespects. The sprending of this and the following of fourth coat w.c. in situ and under their eyes; hence they alone are to blance for any errors in this. After a number of months, the fifth or fine lang coat, composed of the above white lead and zine exide mixture, also a small quantity of French order to make the coat cream codored, was applied.

In a not pretigularly suggest accounts and its tree part, the lead

In a not particularly severe exposure and in two years, the last three lead and zine contings were disintegrated and washed away in large areas over the entire structure, showing the foundation coats of red lead. A cloth wetted in water and wiped over the contings washed them off as freely as though they were of whitewash. Wringing the cloth left the three coats of lead and zine pagments in the water like so much chalk. If the percentage of zine oxide in the coating had been forty, the paint would have failed by preling in strips instead of chalking. The paint, in its materials, proportions, application, location, and exposure, is not far different from that of the New York City elevated railways, which thus far has kept its place free from chalking, but has not presented correction from attacking every foot of the structure beyond the possibility of correction.

Had the same amount and quality of the paint been spread on a thousand inland structures, the failure of the coatings would not have been so marked as in the above case, where the agency of acres of decayed paint tells the tale; but it would have occurred just the same, though the loss would have been so widely distributed as to call no special attention to it.

The reliable qualities of a paint countered of a purcher of

ble nature of that class of pigments.

It is the photograph of a four-inch-diameter wrought-iron pipe. The upper end, A, was enclosed unpainted in a cast-iron ring. The lower end, B, is the adjoining part of the pipe that had been painted with two coats of sublimed lead and zine. The pipe and its connections were buried in the earth for nine years, When taken up the end A had corroded and lost over it inch in diameter. The part B was uncorroded and nearly all of the paint on the whole length of pipe was unchanged and in place. Where the coating was sealed off in the process of removing the pipe from the trench,



Fig. 46,-Four-inch wrought-iron pipe.

the iron was as clean and uncorroded as when laid. (See Chapter V, Sublimed Lead.)

Sir Benjamin Baker has concisely remarked that "it is the deviation from the average which really is so important in the design of engineering works."

This is equally applicable to the design of a paint. The majority of paint-manufacturers appear to harbor the idea of the universality of their product. They give little or no attention to location, exposure, or the many influences to which it may be subjected during its life on a ferric structure.

A paint that is reliable in open inland locations often fails on the seacoast, or in manufacturing towns, or on industrial establishments. The paint that is reliable in the latter place may fail quickly on another not far distant location because of disregarding one or more of the above factors. one of the group; for instance, where red lead or red lead and lampblack are added to influence the drying of the paint. Another instance is where a substance combines in definite proportions, such as the carbonate of lime added to an iron oxide to neutralize the sulphur in the pigment, or the acid in the vehicle.

It is also to be kept in mind that every metal is electropositive to its own oxide; the latter induces corrosion in the former whenever the two are brought into contact, as in a paint conting. The vehicle only insulates or protects either substance but indifferently, particularly where the covered metal is the base of the oxide in the paint. The water and acids in the oil (the latter not infrequently rancid) have great influence in the decay of the conting and corrosion of the covered metal.

The production of enamel paints has become a tracle of great importance. They differ from ordinary oil-vehicle paints inasmuch as they dry with a high lastre or gloss. They were first called "varnish paints," for the reason that the vehicle was a varnish, instead of linseed-oil. They are used principally for conting walls on the inside of buildings that require to be washed with water, as in hospitals, courtyards, etc., the varnish vehicle being less easily injured by the water than an oil vehicle.

The use of a varnish vehicle for ferric paints has been tried in a number of instances in late years with varying degrees of success. The principal difficulty in their use is the uncertain character of the varnish vehicle, which requires a greater knowledge of the nature of the fossil resins and how to compound them than the average paint-manufacturer has at his command.

There are three classes of enamels: *

First. The slow-drying enancels, that require from twelve to fifteen hours to dry under normal conditions. They give a fine, lustrons, level conting, and if carefully applied should show no brushmarks. They are essentially an oil and fossil-resin varnish, with which the necessary color pigments are ground.

Second. Quick-drying enamels, that dry in from twenty to forty minutes according to their composition. They are essentially spirit varnishes, colored with pigments. They have a histocless or flat

Third. Baking enamels. They are varuish compounds that when heated flow into a uniform and lustrous coating. The pigments added to them give them body and color. Sewing-machine and bicycle frames, hurdware, etc., are examples of these coatings. In larger ferric bodies they are represented by the baked japans applied to water-pipes. (See Chapter XI.)

The composition of the varnish base of all of these enamels is the essential part, and in most cases better results can be attained if the varnish is obtained from a reliable varnish-manufacturer than where the painter makes it himself. Even the best of varnish-makers fail sometimes to produce a reliable varnish owing to many causes, principally from the use of common resin and other poorquality gams, the effect of which is the "cruzing" of the coating.

The general nature of the varnish base is indicated in the following recipes that have given good results:

Sixty pounds of good, white Sierra Leone copal, mixed with 10 gallons of the best quality of hot boiled linseed-oil. When well cooked add 16 gallons of turpentine and ½ pound of linolente of manganese for a drier. This varnish, suitable for all colors from black to white, should be ground in as for oil paints.

A cheaper grade of varnish can be made from 35 pounds of Kauri gum and 15 pounds of Sierra Leone copal, mixed with 7 gallons of hot boiled linsced-oil and 1½ gallons of turpentine. When nearly cold, thin down by adding 10 gallons of turpentine. This will be too durk for white enamels, but answers for all other colors.

The white enamels are usually made by adding zine-white or lithopone to the varnish. They work well. About 6 pounds of the white is required to a gallon of the varnish. Whiting and pipe-clay are detrimental; they make the conting gray, are deficient in body, and are liable to cause "peeling."

Black enamels require 4 pounds of lampblack to about 6 gallons of either grade of the above varnishes.

A quick-drying varnish base is made from Sandarac gum, 10 pounds; soft Manila copal, 5 pounds; gum benzoin, 1 pound; methylated spirit, 8 gallons; or 4 gallons each of methylated spirit

drying varnish is made from gum dandour, 14 pounds, and 3 gallons of turpentine.

For a white enamel from either of these varnishes grind in 10

For a white enamel from either of these variables grind in 10 pounds of zine-white or lithopone to 1 gallon of varoush.

For a black enamel grind in 5 pounds of zine white, 2 pounds of carbon black, and 3 ounces of brilliant chony spirit black to 4 gallon of varnish.

The baking enamels are not essentially different from the first class of variashes herein mentioned, other than that they contain more gum or resin of some quality.

A few special recipes are the following: Best grade of refined asphaltam, 70 paniels, mixed with 9 gallons of hot lineseed oil and 5 gallons of gold size. Boil until ropy, then add 9 gallons of turpentine. This is for a black conting.

The colored enamels require a better grade of varnish base, which is made thus: A good quality of Kanri and copal games, each 20 pounds; animi, 5 paunds; melt together and mix with 14 gallons of hot linseed-ad; boil until stringy, thin with 18 gallons of turpentine, and use with any pigments to get the desired color.

A quick-drying black cunnel is composed of:

D. C. shellac	141	รูพหมากโซ	
Gam sandarac		• •	
Gum benzoin,	* \$	25	
Latopblack		11	
Custor-oil,		t i t	
Spicit aniliae black			
Methylated spirit			25 gallens
Wood-spirit			25 11
Total	ни	25 pannds	27 galleon

The pigments assembled with all of the above varnishes should be of the best quality, particularly the lampblack. Pulverized bituminous real and nearly all of the so-called "carbon blacks" prove detrimental to the quality and color of the enamel. No benzine or turpentine can be added to any of the above varnishes after they have left the cooking-kettle without affecting the gloss of the tective value of all enamel paints is quite as variable as the ordinary oil-vehicle paints. A varnish vehicle does not always secure a reliable ferric coating, unless more care is exercised in preparing the surface to receive it, or in its application, than the average painter generally gives to these matters. The use of resin or resin-oils in enamel paints is as disastrous as when used in an oil paint.

CHAPTER XXXIII.

CORROSION OF IRON AND STILLS

THE difference in the rate of corrosion between iron and steel as given by different authorities varies greatly. The reason for this is plain; the conditions of each reported rate of corrosion, whether the result of laboratory or other tests, such as exposure to weather or other corroding influences, not being similar in all respects, are therefore comparable only in a general way.

Pieces of iron and steel, both suitable for boiler-tubes, were made clean and bright, then placed in sandy loans with which had been thoroughly mixed some sodium carbonate, sodium nitrate, ammonium and magnesium chlorides. The earth so prepared was kept moist. At the end of twenty-three days the plates were taken out, cleaned and weighed. The following was the result:

fron had	OSL	by	corrosiou.	 cent.
Steel "	61	"	4	 * *

The pieces were replaced in the earth and left for twenty-eight days longer, or sixty-one days in all. The result was:

Iron	total	loss	by	corrosio	11	2.(Hi)	H-1.	cent.
Steel	"	"	u	**		1.79		

This is a rate of corrosion that would probably have caused the disappearance of the plates inside of eight years

Experiments conducted by the Admiralty, Board of Trade, and Lloyds prove that steel corrodes much more rapidly than iron when exposed to the action of salt water; also that the conductor brands of iron corrode less rapidly than the better brands when exposed to the same influences. With steel and iron both unprotected and exposed to the same action of the weather and sea water corrosion advanced at the rate of one inch in depth in \$2 years for the steel and 190 years for the iron. When always immersed in sea-water the periods are one inch in 130 years for the steel and 210

by four inches would be entirely corroded away in a little over 100 years.

Mr. G. Rennie's experiments in 1836 were with embes of wrought iron, cust iron, and brouze for lighthouse purposes. The cubes in separate vessels were impressed for seventy hours in saline solutions considerably stronger than sen-water. The cast iron lost $_{33}^{1}_{03}$ of its weight; the wrought iron $_{6.6}^{1}_{10}$, being in the proportions of two to one in favor of cust iron. The brouze lost $_{10.600}^{1}_{0.0}$ of its weight, a result in favor of brouze over cast iron of three to one. The east- and wrought-iron cubes were then placed in a strong solution of one part of muriatic acid in twenty-five parts of Thames water, and exposed for twenty-one hours. The east-iron cube lost $_{33}^{1}$ of its weight, the wrought-iron only $_{3.68}^{1}$, being eight to one in favor of the wrought iron.

In these experiments with the same samples of each metal the results were directly contrary. The crystalline nature of the east iron evidently favored the disruption of the crystals from their bond or loose association together; they were east off when partially corroded, and corroded by themselves while the acid had fresh surfaces to act upon in all directions.

The experiments of Mr. Robert Mallett, M.I.C.E., on wrought iron and east iron sunk in the sen, showed that from τ_0^1 to τ_0^4 inch in custings one inch thick, and about τ_0^0 inch of wrought iron, will be destroyed in a century in clear sen-water. This is equal to fifteen to one in favor of cast-iron. Other experiments by Mr. Mallett showed that cust iron unprotected and exposed freely to atmospheric action was corroded nearly as rapidly as by the action of clear seawater.

Mr. Kenniple, Central India Railway Company, reports that "the greatest corrosion of east-iron piles existed close to the low-water mark, and did not extend to any considerable distance from that point." This condition he also found to exist in the wrought-iron bolts and braces. After an exposure of twenty-five years the piles were found to be in very good condition, and corrosion had only occurred in places accessible for renewals. A thin coating of mud, marine growth, and barnacles upon the immersed surfaces of the

once. He concludes that after a life of from thirty to bity years east-iron structures exposed to seaswater can only be regarded as of a temporary character, especially those of light east from pile design.

Dr. Grace Calvert, F.R.S.,* experimented on a number of gray cast-iron cubes made of Staffordsbire cold blast from fromersed in acidulated water. The specimens were 0.39 inch cube, specific gravity 7.858, weight of cube 237 grains. The cubes were placed separately in bottles holding about 30 cubic inches of greatly diluted sulphuric acid. Similar cubes were placed in bottles containing dilute hydrochloric, acetic, and phosphoric acids. The action of the acids on the iron was slow; but at the end of three months, although the appearance of the cubes had not changed, some of them, especially those immersed in the solution of acetic acid, had softened so that a knife-blade could penetrate them 0.11 to 0.16 of an inch.

The solutions of acids mentioned were replaced by a fresh one in each bottle every two months for two years. Changes were then found to have taken place in all of the cubes, the acetic acid showing the greatest decomposing effect, then the hydrochloric, sulphone, and phosphoric acids; the latter had the least effect upon the cubes,

The action of the acids upon the iron had changed its nature, without any alteration of its bulk or in the appearance of the preface of the cubes. The weight of one cube after two years' immersion was 54 grains against its original weight of 237 grains. Its specific gravity was 2.751 instead of 7.858. The change in the physical character of the iron is indicated by the following analysis of the gray east-iron from which the cubics were made, and a set of cubes after a two years' immersion in the acetic acid solution:

	Refu	ir lii	tistet ansst	Afri	nt 1::::::::::::::::::::::::	
Iron	. 95	113	pet cent	711	fifill ju t x e tel	i :
Carbon, . ,		HIII	٠.,	11	070 "	1
Nitrogeo	()	7111	**	2	THE "	į
Silicon,	- (1	178	**	- 6	11711	- 1
Sulphur	0	179	**	(1	ten, "	-
I Phosphorus,	()	132	**	0	059 **	- 1
Loss,	0	108	**	1	155 "	ì
	-			1	• • • • •	ī
	100	OOO		1/81	Luus	1

of the atmosphere, and rating them as 0, in Glasgow they were 83, and in London 28.

The comparative amounts of ammonia and other impurities in the air and in min-water were: Valentia 1, Glasgow 50, Liverpool 30, Manchester 36. For the amount of hydrochloric neid present in the same elements Blackpool was 100, London 320, the Underground Railway in London 974. Anhydrons sulphuric acid at Blackpool 100, London 352, Underground Railway 1554. Ammonia and albuminoid ammonia at Innellan, on Firth of Clyde, 100, London 108 and 117, Glasgow 150 and 221, the Underground Railway 138 and 271.

Drs. Clowes and Andrews' examination of the air in the cars of the Central London Railway showed a maximum amount of earbon dioxide of 44.7 volumes and a minimum amount of 9.6 volumes in 10,000 volumes of air. In a railway-station elevator 15.2 volumes was found.

Many points in the Paris underground railway system have 33 volumes of carbon dioxide in addition to other deleterious gases and 2 per cent of aqueous vapor. Dr. Clowes states that not more than 8 volumes of carbon dioxide in 10,000 volumes of air should ever be present at any point in a railway building.

Dr. Smith also found a variety of solid substances in the air, such as common salt, sulphur, nitrate of ammonia, lime salts, iron; also the phosphates, iodides, and other organic matters given off by animals, vegetables, etc. The percentage of oxygen in the open air varied from 21.0 to 20.40, while the carbonic gas varied from its normal amount of three parts in ten thousand to over thirteen parts in the London Underground Railway, and some parts of the Swiss tunnel contained over 17 per cent.

Dr. Huxley's Physiography gives his examination of the amount of carbonic acid gas in 10,000 parts of the atmosphere at a number of points as examined by him, viz.:

and soot in the air, in the central district of training in the year 1902. The fall of dust and soot in an open dish of 75 square inches area amounted to 2 onners, equal to 3.8 oracles per square foot, or 23.5 pounds for every 100 square feet.

The above amounts of corrosive elements are not arbitrarily constant, but they indicate the corrosive influences that may be encountered in almost every situation of engineering work.

Mr. Beardmore, C.E., instances a case of a sea lock, in which soft water was "locked" down into the sea water level. At the end of thirty-five years' service all of the cast and wrought non attachments to the wooden lock-gates, also the spike in the platforms and gate-sills, had completely corroded away, though the timber pairs of the structure were perfectly sound. Ferric metal exposed to the action of salt or freely water which

is not changed corrodes far less rapidly than where the water is changed more or less frequently.

changed more or less frequently.

Dr. Lyon of India, a high chemical authority, reports that some east-iron piles, after four and a half years' expective to the action of pure sen-water having a specific gravity of 1 028 and that

contained 3000 grains of solid matter per gallon, of which 1605 con-

sisted of the chlorides of sodium, magnesium, etc., had undergone a change to a depth of τ_0^1 inch from the surface of the metal.

The Milton-on-Thames pier was erected in 1844 on cast iron columns 3 feet in diameter and 15 inches thick. In the Grave end town pier and the Maplin Sands lighthouse the cast iron columns and other castsiron members exposed to the sea water were originally 14 and 14 inches thick. In all of these structures, at the end of texts five years, only 3 inch of the metal rem, incd unaffected, the rest had changed to the semblance of plumbage. None of the members

of these structures indicated any change in the metal.

Mr. Thomas Rhodes, C.E., reports that "in the lock of the Caledonian Canal the cust-iron sluice-gates were expected to carwater. All of the parts were control with a heavy Swistish tar, except on the working faces of the gates. These faces were faced

and ground together. Four years after the immersion of the sates, upon an inspection of their condition, on all of the parts coated with

of § of an inch, and had to be renewed."

The experience of American engineers appears to be equally conclusive of the treacherous character of east iron exposed to sea-air or sen-water.

Mr. John D. Van Buren, in a paper read before the American Society of Civil Engineers, stated that "bolts and other wrought-iron parts are badly corroded in less than twenty-five years when submerged in sea-water. Certain kinds of east iron could perhaps be made to last fifty years, which would be a generous allowance, and probably greatly exceeds the average life of east iron exposed to sea-water."

Sir Benjamin Baker, in his paper "The Metropolitan and Metropolitan District Railways," * snys: "In tunnel constructions when the roof members and bottom flange of the girders, tie-rods, anchors, etc., are much exposed to corrosive influences, wrought-iron members were used and thought to be more trustworthy than east iron, but were found to be exposed to a greater risk from hidden oxidation. Experience has shown the trouble and cost of maintaining ironwork exposed to atmospheric corrosion in an underground railway. It is so great that it would justify a considerable increase in the first cost by substituting brickwork and deep cuttings for ironwork and shallower construction. Where the depth was sufficient for an arch, brick covered ways were to be preferred to iron-girder constructions, on account of the smaller cost, increased durability, and safety,"

Sir John Fowler confirmed Sir Benjamin Baker's views in regard to the substitution of brick work and musomy for ironwork in all cases where possible, even at a material increase in the cost of the work. He thought the question was not confined to the relative rate of corrosion of east iron, wrought iron, or steel, but to the great risk arising from hidden oxidation on important members of walled-in ironwork.

These views are corroborated in the experience of American railways. The Pennsylvania Railway has already replaced a number of its iron bridges with masonry arches, and wherever possible will continue the substitution.

Car-wheels made from the best of gray cast iron when immersed either constantly or alternately wet and dry, in fresh or salt water in all stages of impurity are found to corrode faster in the body part of the in both parts of the wheel and under the same corresive influences. The difference must be attributed to the effect of the chill, which has changed the molecular formation of that part of the wheel, making it more dense and of a needle like or filamentous formation, that is not so readily attacked by corrosion as the crystalline part. This effect in car-wheels is the more noticeable as the body part of the wheel is covered with a skin of the silicate protoxide of iron, produced by the molten metal fusing the sand in the mold. There is also a further protection afforded by a film of the black magnetic oxide of iron being formed at the same time by the oxidation of the hot metal in cooling.

Cast-iron pipe or piles would be more durable were they cast from chills the same as the trend part of a car wheel. Instead, however, of this method, which is attended with some difficulties not present in the casting of car-wheels, it has been proposed to increase the depth of the silicate coating on such bodies by a process used by the ate Mr. E. F. C. Davis, late President of the American Society of Mechanical Engineers, in the protection of mining pumps, chambers, and pipes subject to the action of mine-water. The corresive action of this fluid is very great and increases by notion and pressure. The process consists of coating the cores and other parts of the mould with a thin paste applied by a brush, which increases the thickness of the silicate coat to more than double its natural thickness, and can be made still thicker by repeated applications of the paste before casting. The composition of the paste is, viz.

112 parts silica.

44 " enleined soda earlemate

24 " " enthante precipitate

4 " borneic neid.

Mix and thoroughly pulverize them. Cost the core or mould with plumbago facing, and apply the enamel as a powder or parte to the thickness required, in one or more applications, and east as usual. A single coating with this compound more than shouldes the life of the metal.

inside of five years changed to a plumbago-like substance, that had hardly any strength and could be cut easily. No change in the appearance of the metal was indicated; the change in the metal was from the inside, or where in contact with the water. The pipes had the usual tough skin, and were nominally protected by a foundry dip coating. Mr. F. A. Boyer, M.E.,* reports that a cast-iron water-pipe 12 inches

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in dimmeter, 1200 feet long, used to circulate sea-water for condensation purposes, laid mostly in the open air, was changed in two and one-half years to a plumhago-like substance that could be cut easily, but to the eye presented no indications of the change in the metal.

Gray east-iron water-pipes \dagger laid at Atlantic City, N. J., for nineteen years, in black swamp-mud containing decomposed vegetable and saline substances highly acidulous and corrosive, were found to be generally corroded externally. The iron was softened about $\frac{1}{k}$ inch in depth, and in many places it had extended through the pipe. The corroded metal appeared to have been replaced by a clay-like substance of light weight, containing 17 per cent of silica, particles of the cast iron being embedded in it.

A member of the American Water-works Association reported in the 1902 meeting the decay of two cast-iron suction-pipes, where the metal was changed to a graphitic substance, easily cut. The suction water was originally soft ground-water, but later was artesian well-water containing a small amount of sulphuretted hydrogen. Wherever brass and iron came in contact with the water-mains, eventually the iron became soft enough to cut easily.

Mr. Trautwine, an eminent American civil engineer, recommends a white close-grained east iron or chilled iron for piles and wharfs exposed to sen-water or sea air.

Mr. Turner, Assoc. R.S.M., recommneds a gray cast-iron for docks and piles exposed in English sea locations.

The above results of the use of different brands of cast iron for sea-air and sea-water exposures in different locations are probably due to the difference in sea-water in different parts of the world. Sea-water varies greatly in corrosive and fouling properties even if not contaminated as mentioned before.

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Cannons and other tough cast-iron articles, 6 or more inches in thickness of metal, sunk in the sen in many parts of the world, change in about one hundred years to a soft carburet of iron or a plumbago-like substance, without any diminution in size or any exfoliation in scales or flakes, as in atmospheric exidation. They become too hot to handle, when first raised and exposed to the air, from the absorption of oxygen.

Mr. Francis T. Bowles, Naval Constructor United States Navy, states: "That the corrosion of fron and steel, from the observations made at the different Government navy yards, with the different kinds of iron and steel used in inval vessels exposed to sea water in various parts of the world under a great number of conditions of temperature, brackish, sewage, and dock water, was: That unpainted iron and steel plates will carrolle in one hundred years on each exposed surface 30 to 50 inch of metal; in ordinary fresh water, 02 to .03 inch, and in the atmosphere, .25 to .30 inch." But he makes no distinction in the rate of corrosion between iron and soft steel.

Mr. L. M. Hastings, City Engineer, Cambridge, Mass., reporting the results of his experiments to ascertain the difference in corresion between wrought iron, soft steel, and east non, all uncoated, exposed to a running mixture of pand and brook water, tairly soft and comparatively free from any acid or saline matters, states: "That after an exposure of one year the uncoated wrought iron was badly tuberculated and rusted, the soft steel was similarly affected, but not to the same degree; the east iron was also affected. The percentages of increase in weight due to corresion were as follows. The wrought iron, 1.5 to 3.1 per cent; the steel, 1.1 to 2.4 per cent, and the east iron, 1.0 to 0.96 per cent. A similar set of plates barred in sand, also in clay soil, showed the same relative difference in corrosion of the metals."

Dr. Robert II. Thurston* reports the result of his investigations of the different rates of corrosion between iron and steel. Directly, they are: "Cast iron in dilute solutions of neids is rapidly acted mon, the matal requiring its ground account.

rusts less rapidly than soft steel. Bilge-water corrodes iron and steel rapidly. Iron ships carefully painted have been found to corrode at a rate not far from ι_{0}^{1} inch in twenty-five years."

Thwaite* gives a formula and table of constants for the rate of corrosion between different metals under different elemental exposures, all for unprotected metal, viz.:

		Wu	ter.			
Matorial.	Hon.		River.		Impure Air.	
1	Foul.	Clour.	Foul.	Clear Water, or in Air.		
Cast iron, Wrought iron Steel, Cast iron, no skin, Calvanized iron	0,0056 ,1956 ,1944 ,23 ,00	0,0636 ,1255 ,0970 ,0880 ,0359	0.0381 .1440 .1133 .0728 .0371	0,0113 .0123 .0125 .0100 .0048	0.0476 .1254 .1252 .0854 .0190	

Average for sen-water: Cust iron, in contact with brass, copper, or gunbronzes, 0.10 to 0.35; wrought iron, in contact with the same, 0.3 to 0.45.

No analysis of the constituents of the several metals is given, and the terms hard and soft metal are very variable conditions. However, all of the above experimenters give hard metal, whether of cast iron, wrought iron, or steel, as being less corrosive than soft metal.

Mr. Thomas Andrews, F.R.S., experimented at the Wortley Iron Works on wrought-iron and steel plates containing varying amounts of earbon. The plates were immersed in sea-water that was changed monthly. It was found that the lower the percentage of combined carbon in the metal, the lower was the corrosion. The best wrought iron corroded less than any of the steels at any stage of their exposure during the one hundred and ten weeks of the test. Wrought iron that contained double the usual amount of phosphorus and

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In any of the steels, manganese in excess tended to produce an increased corrosion, evidently from its unequal distribution and the galvanic action on the adjacent metal.

On the whole, from the many reported cases of corrosion in all parts of the world, that include many quadries of eact non and wrought iron that have had approximate exposure, for thirty or more years, it appears that hard close grained and chilled iron are less liable to corrode than any brands of softer metal. Pessenier, open-hearth steels, also steel castings used for structural work, have not yet had time enough to afford many comparisons with each other, or with east iron and wrought iron, to prove which metal is the most affected by corrosion.

It is left to the future to develop some alloy of non-or steel that will retard if not prevent corresion, while not materially reducing their strength or other qualities.

The composition of wrought iron and the processes it is subjected to between the bloom and finished article have a great effect to determine its rate of corrosion. Iron containing sulphur is ted short, that containing phosphorus is cold short. Both differ in corresphility; the cold short is the one less affected, being harder and more crystalline in composition, while the red short has the sulphur element to aid corrosion. Neutral iron made from both of the above brands has a different rate of corrosion than either

The same quality of iron worked in the rolls, in the one case both lengthwise and crosswise, to produce sheet or plate iron, differs in corrosibility from that worked principally in one direction, as in the case of beams, angles, and other structural shapes. All of the latter forms tend to disintegrate by corrosion into trips, needle or fibrous form, owing to the granular character of the iron being changed by the rolls into parallel fibres, that are not interlocked as the cross-rolling arranges them. The corresion aided by the cinder follows the grain of the metal. The cinder is acid and porous, and only in mechanical bond with the iron by reason of the action of the rolls.

through the rolls. Enough, in fact, to show a marked difference in the corrosion of the two products, when from the same metal. Boiler-tube skelps, being made from a better quality of iron, or having been refined by a further working of the same grade of metal, by having had some cross-rolling before being made into skelps, are less affected by corrosion than ordinary wrought-iron pipes. The arger sizes of these pipes being made from long rolled skelps and hip-welded, instead of only welded, as in the case of small pipes, show in the welds a less corrosibility than in the body part of the pipe. This is

due to the additional condensation of



Fig. 47. Laminated corrosion of a steel-plate girder on the Washington Street railway bridge in Boston.

the fibres at the welds; also there is less einder in the lap-weld.

Cold-rolled shafting and rods are rendered more dense by the rolling process, and they are less affected by corrosion than the bars from which they are cold-rolled. The process also increases their tensile strength. Cold-drawn wire also presents the same features. A wire nail corrodes less than a cut nail, so does a hammered or so-called wrought nail.

Polished iron and steel tools, sword-blades, razors, etc., resist corrosion better than the same articles ground, but not polished; the improved resistance being due to the surface not having so many small cavities to hold any moisture reaching it, because of the condensed and repellent nature of the surface due to the polishing.

Burnished surfaces resist corrosion and are more repellent of moisture than polished ones; but, if corrosion is once established on them as a spot, it appears to concentrate an energy to produce a deep corrosion, that is difficult to eradicate.

Rivets have a different rate of corrosion between their heads, or points, than the body of the rivet. Corrosion of a riveted joint

The quantity of metallic iron in the best refined brands is 99.8 per cent. In common bar iron it is 98 to 98.3 per cent. In commary east iron, about 93.5 per cent, and 2.5 per cent of graphatic carbon,

The percent; of volume in ordinary east iron is 1.41 per cent; in different kinds of Bessemer steel, 0.41 to 1.20 per cent. In a locomotive (ire ingot, 0.57 per cent. For a hard iron tire, 0.97 per cent. In a basic-steel ingot, 1.22 to 2.17 per cent.

At an annual convention of the American Institute of Architects, in a discussion on the use of iron and steel in the construction of modern high buildings, it was reported by one of the leading architects in the United States that the iron beams removed by him from the old Times building, though in use only thirty-five years, were rotten with rust. They were enclosed in eight inches of brickwork, forming the arches that supported the pavement over the vault where the steam-boilers were placed, and though always dry, yet had been exposed to ordinary fire-room vapors. They had been well painted with iron-oxide paints and protected from external moisture by an asphalt covering. The iron came off in strips, clearly showing that the rust had followed the landination of the iron, the web of the girders being so rotten as to be easily broken by the fagers. Other examples of sidewalk beam corrosion are given on page 260.

Anchor-stocks made from hammered iron always show here corression than cable links. In both cases only the best quality of iron, that contains but a small quantity of einder, is employed. The link corrosion is in the form of strips, following the fibre of the metal, while the anchor-stock generally corrosies in a compact scale form. A different rate of corrosion exists in the cable links at the end, where they are welded from that shown in the body. The iron in or near the weld is refined and more dense, also has the fibres interlaced; all these points have a tendency to delay corrosion, which is more rapid where the fibres are undisturbed by the hammer.

Tunnel Shields and Submarine Metal Corresion.

With the recorded instances of the corresion of iron and steel to judge from, it may be pertinent to ask, how love will the contact living

when completed, these constructions have a small margin of strength above that necessary to get the metal into place during construction, and none at all as a reserve for the loss in strength from the inevitable change in the metallic work exposed to sea-water, which begins just as soon as the shield is in place; and in the case of tough



Fig. 48.—Corrosion of a steel plate from the Washington Street railway bridge in Boston.

close-grained cannon metal, has been ascertained to be at a rate of about six inches in one hundred years.

The lining plates and ribs for tunnel-shields are seldom over three inches thick. The different sections of the shield utterly preclude any material strength to be derived from their circular form, made with bolted joints. The bolts holding the shield sections together and to each other are relatively small, and will be the first to yield to the effects of saline corrosion.

The brickwork or concrete lining of the tunnel, however thick, or the thin coat of partly dried paint, will not protect the shield metal, for any appreciable time, from the change to a plumbago-like substance, which does not require the presence of air to produce it. The passage of railway trains through the tunnel will set up an undulating or ribratory movement through the tube resting on its hed

However strong such metallic shields and maintry lined constructions are when driven through took or earth, in or through salt water or the saturated six of salt water, they are the most treacherons and dangerous of all engineering devices yet conceived. affecting the transportation and satety of the public

Metallic salts and acids in water intensity the corresion of all metals exposed to their action either by maneraton or by combination of the vapors from them. The metal work of radway tunnels is disastronsly affected by the condensed vapors of sulphanous and carbonic acid and the moisture due to such legations, the corresion of the metals decreasing the resistance of the water to voltaic circuits; this corrosion by liquids being voltaic phenomena in all cases, and in many cases is intensified by the moisture being in the form of drops instead of being uniformly spread over the whole surface. The cut (Fig. 40) from the Radroad Gazette, November 23,

1894, represents a section of a seventy six pound tertail hid the Museometeoric Tunnel, removed after being five years, having lost more weight by corresion than wear, The dotted lines show the original size of the rail, and the full lines its present worn and corrolled size, which is very marked. The rails were removed on account of their strength having been seriously affected by the corrosion. The tunnel is very damp, and a great deal of sand is used by the engines, which kept the base of the rail covered, the vibration caused by the passage of the trains having a tendency to remove the thin scale of rust almost as rapidly as it could be formed. There was but little apparent difference in the corrosion, whether between the cross-ties or where the

Pag. 49.

almost motionless for twelve hours per day, and though the accumulation of carbonic acid is rapid, and a part of it is absorbed by the grent quantity of water present, the air is almost unrespirable, and causes a great deal of distress to the workmen, and the corrosion of all metal-work inside the tunnel is very rapid.

The water that trickles down the walls of the tunnel is the condensation of the exhaust gases from the locomotives. It contains suphuretted hydrogen, sulphur dioxide, annouia, and earbon dioxide, and other combustion gases. The rails are renewed every ten years, and the telegraph cables in the tunnel require exceptionally strong wrappings.

In the new Simplon Tunnel, 64,718 feet long, forced draft is proposed, requiring over 500 H. P. at the ventilator shaft. The fans render an effective duty of 65 per cent, 1760 cubic feet of air per second being required for ventilation.

In general, the corrosion of metals in tunnels where the rails are bedded on cast-iron chairs is represented by east iron, 100; wrought iron, 120; steel, 133.

In the Arlberg Tunnel, 33,587 feet long, the corrosion of the rails and other metals is so rapid, that they all require renewal every ten years. The corrosion comes principally from the condensed gases of the locomotives, though the traffic is light and a good quality of coal is used. The temperature of the tunnel remains almost uniformly 75° F. throughout the year.

The amount of free sulphuric acid in the exhaust gases from tunnel locomotives using a good quality of bituminous coal has been found to be about 5 pounds per hour, varying from -0.3 to +7.9 per cent.

In a tunnel in France 2850 feet long, where but little water came through the walls, the 78-pound-per-yard rails were replaced after the passage of 230,000 trains at a speed of nineteen miles per hour. They had been eleven and one-half years in service and had lost in weight 184 pounds, or 24,166 per cent per yard. The corrosion was general over their whole surface, but the rolling action of the wheels on the head increased the corrosion at that point by keeping the motal bright and removing the rust as fast as it formed

This coating was firm and tough, and did not theke or scale off in transportation or in the laying of the carls. At the end of twenty years the metal had not corroded to any appropriate extent, except where mechanically injured. This conting was practically the conttar dip used on water-pipes by English tounders.

Mr. Otto Herting cites the instance of the corresion of some tee-rails used as girders in a Cape Breton ione, that had been abandoned twenty years. The metal was changed to a gravish brown color, could be cut, and had a specific gravity of only 2.053. The metal powdered in a mortar was magnetic. It analyzed as follows:

Iron	31 30 per cent.
Graphitic carbon	21 10 **
Silicon	11/20 ***
Manganese	1 93
Sulphur	1 (8)
Phosphorus	à 85 · · ·
Undetermined and hose, .	21 12 **

Ithi thi .

Fig. 50 represents the corresion of steel tails land upon docks and other places contiguous to sea water, where the effect of corresion was equal to about 4 per

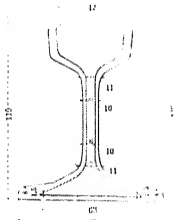


Fig. 50.—Rail section of the Sumatra Railway, showing the

cent each year upon the weight of a 32 pound rail per vard. The details of the rail from which the cut was tunde were contributed by Mr Delpint, Chief Engineer of the Sumatra State Railway, through Mr J. W. Post, Divisional Chief Engineer of the Netherlands State Railway.

The annual report for 1980 * of the Samarang Joana Steam Transway Company (of Java) states that a considerable number of steel rails had to be removed from the harbor tracks on account of corrosion, which amounted to 10.7 kilos sifying the action.

Thermo-electric currents arise from changes in temperature and set up voltaic action which, though slight and not casily detected, will colarge all fissures, cavities, and seams sufficient to sap the

from decomposed pyrties and other innerals, aided by near, inten-

strength of the metal.

Dr. Henry Wurtz* proposes an electro-chemical process for protecting mine metal, by connecting all of the fixed metal in the mining plant as the negative element, with a dynamo of sufficient force to overcome the galvanic energy of the surfaces when exposed to the mine's corrosive liquids, the positive terminal to be connected to a mass of hard coke in the mine sump. These conditions vary but slightly from those existing in a ship, and it is not improbable that both systems could be made to work effectively.

In a number of instances where the whole system of mining-pipes required renewal every two years, corrosion was completely stopped by conting them with an enamel in the following manner:

The pipes were first pickled in a bath of hydrochloric-acid solu-

tion to free them from the foundry scale, then washed thoroughly, and dried. The pipes then received a coating of 34 parts of silica, 2 parts of soda, and 15 parts of borax. These were mixed in a little water and the pipes exposed for ten or fifteen minutes in a dull red-hot retort. A second coating was applied, composed of 34 parts of feldspar, 19 parts of silica, 24 parts of borax, 16 parts of oxide of tin, 4 parts of fluor spar, 9 parts of soda, and 3 parts of saltpetre. These were melted together in a crucible. When cold the mass was ground to a fine paste in a little water and applied to the pipes with a brush. The pipes were then exposed in a muffle to a white heat. The enamel so formed thoroughly united with the iron, and has protected the pipes for over forty years, and they are apparently in a good condition now.

The Journal of the Society of Chemical Industry (London), February 28, 1894, details some experiments upon the galvanic action of sen-water upon iron and steel structures in various relations with each other, as constructive parts of trusses, boilers, etc., to prevent the corrosion. The use of zinc and other easily axidized metals and allows is corrected to the second allows is corrected to the

the structure, or at least aid the paint conting in its interior of protection.

These protective features, proposed for the internal parts of a ship, do not apply to the protection of the external surfaces, where an entirely new set of conditions are in toric, owing to the numerous rivets employed to hold the plates together and to the traines, and which are necessarily improtected from the many sources of corresion herein mentioned.

Professor V. B. Lewes* of the Royal Naval College, Greenwich, England, at a recent meeting of the Institute of Naval Architects, London, states:

"The rusting of iron and steel is a definite chemical process, due to the conjoint action of air, mosture, and carbon dioxide upon the metal. The increased rate of chemical corrosive action due to a local increase of temperature is noticeable, and may be due to galvanic action set up between portions of the same metal at different temperatures.

"It is an undoubted fact that the double bettern of ship plates near the boilers corrodes more rapidly than smallar plates in other parts of the vessel, and the increase in temperature near the boiler is the only factor.

"It is also noteworthy that the plates at the bottom of the cellular spaces which are kept cool by contact with the sea water do not corrode; and cases are noted in which parts of a plate, which get locally warmer than other parts—although the difference can only be a few degrees—corrode much more rapidly than the cooler portions

"Experiments show that the rapid corresion found in the double bottoms near the boilers or other sources of heat, is due to galvanic action, and not to the increased chemical activity due simply to the increase of temperature. As the aslas are drawn and quenched with senswater near these exposed plates, no doubt some of the corresion can be traced to the gases thus formed; the sulphur in the aslas also contributing its effect."

Mr. William Thomson, F.R.S., read a paper on "The Influence

^{*} A paper read at the thirtieth session of the Institution of Nacot Architects

of Some Chemical Agents in Producing Injury to Iron and Steel," before the Manchester Association of Engineers, November 25, 1893, in which he refers to the interesting and exhaustive experiments made by Mr. Thomas Andrews, F.R.S., on the galvanic action which takes place between iron and steel, and between iron of different kinds and steel of different kinds, viz.:

"The galvanic action between wrought iron, cast metals, and various steels during long exposures in sea-water." Institute of Civil Engineers, Vol. 1883-84, Part III.

"Corrosion of metals during long exposures in sea-water." Institute of Civil Engineers, Vol. LXXXII, 1884-85, Part IV.

"The relative electro-chemical positions of wrought iron, steel, cast metals, etc., in sea-water and other solutions." Royal Society of Edinburgh, Vols. 1883–1889.

Mr. David Phillips's paper. Institute of Marine Engineers, 1890.

In the above-named articles Mr. Andrews shows that while some varieties of iron and steel remain constantly electro-positive or electronegative to each other, others change, taking opposite positions toward each other, while others again change positions constantly during long periods, these changes always producing rust,

"It can be easily understood that while there is no material voltaic action between two pieces of steel or two pieces of iron, or of pieces of steel and iron, there may be conditions on the surface of one plate or rivet which may act strongly as an electro-negative element, and produce rusting on the metal in contact with it. A piece of iron immersed in weak nitric acid begins to dissolve at once. A similar piece placed in strong nitric acid, touching it for a few minutes with a piece of platinum wire, and then putting it into the weak nitric acid, will not dissolve, it having been rendered passive; and similarly, there is reason why one piece of iron may act electro-negatively toward another piece of the same metal, on account of some slight alteration of its physical properties, by hammering, such as closing the riveted seams of plates, calking seams, setting tubes,

Government to investigate the cause of the failure of the Tay Bridge, reported that where east non and wrought non-were connected by rivets in many parts of the same structure as they were in this one), the rivets and connecting wrought non-work, where connected to the east-iron members of the structure a charact, flanges, spandrels, etc.), had corroded to such an extent as to be below the point of stability by the local galvatur circles borned at nancerous points in the structure where the two metals were in contact, and the corrosion thus established was the cause of the deaster.

Mr. St. John Day, in a paper read before the Institution of Engineers and Shipbuilders, Scotland, I clauary, 1880, stated that "the Ik-inch diameter bolts holding the ties to the pules on the Tay Bridge were so corroded that they would have to be replaced every four to six years. Some of the bolts were found to be corroded away to one-shalf of their original size."

The Scotland Board of Trade now probabits the connection of cast-from columns with wronght from columns or thes

The corrosive action noticed in the riveted sections of the tubular bridge over the St. Lawrence River at Montreal, Canada, referred to before, resembling Fig. 39. Chapter NNTH, was of a smalar nature to the above example—In this case, while the corrosion was of almost unparalleled amount and virility in the whole structure, the rivets that held the floor-beams and track stringers in place, and were under the greatest strain and subject to vibration and shock from the passing of the railway trains, were corrolled the most, though all of these parts were of wrought iron to wrought iron, but varied in quality from common iron to refined iron

An important question presents itself to boder makers: whether it is safe to rivet steel plates to iron plates in steam boders, or even in other constructions, particularly where expected to high tempera-

tion of Civil Engineers (English), Vol. XI.XVII, p. 323, and Vol. I.XXII, p. 284, "The Action of Tidal Streams on Metals." Proceedings Federated Institu-

Society, Vol. XIII, p. 429; Vol. XI.IV, p. 152, Vol. XIIV, p. 176, and Vol. III, p. 114.

"On the Corrodon of Metals in Seg-water." Minutes of Proceedings Institu

other, Mr. Thomson made an extended series of experiments, using a Thomson's tangent galvanometer to measure the electrical currents generated in the corrosion of iron and steel, both singly and in connection with each other, and when impersed in different fluids, viz.: sulphuric acid (one part to nine of water), caustic-potash solution (specific gravity, 1.311), and chloride-of-ammonium solution (specific gravity, 1.033), the latter representing electrically the ordinary concentrated water found in stema-boilers.

The details of the experiments are important, but I will give only the results obtained, viz.:

"When an iron rivet and a piece of the above-mentioned corroded steel furnace plate were placed in contact and immersed in the weak sulphuric-acid bath, at first the steel was electro-negative to the iron, but in a few moments it changed, and afterward the iron was electro-negative to the steel. When placed in the chloride-of-ammonima solution, at first the iron was strongly electro-positive to the steel, and afterward became weakly electro-negative. When placed in the caustic-potash solution, the steel was strongly electro-positive. but the current gradually became weaker and weaker until it practically ceased. A new steel rivet in an iron plate, a steel rivet closed by a machine and held until nearly cold, an iron rivet closed on a mild unworked steel plate, all reacted strongly among themselves. The iron when first brought into voltaic contact with the steel was strongly electro-positive to the steel, being presumably strongly acted upon by the solution, but after a few minutes almost ceased action or became reversed; and so far as the tests demonstrated as a whole, it was to the effect that it was quite as safe to bring iron and steel in close mechanical contact with each other as two different kinds of steel or two kinds of iron. Corrosion was developed in some degree in the contact of all different metals to each other."

It is cited that a number of torpedo boats of the French Navy, that had been constructed within ten years, and that had not made a thousand knots of sea service, were found to be so corroded at the water-line, though well painted from the first with anti-corrosive paints, they had to be condemned for service; while other boats of the same class that had never been in commission, but had been had up under cover, had, as the report says, "eaten their own heads off by corrosion," and were condemned for the same cause. In these cases the corrosion had been in progress under the paint covering, and showed but little sign of its extent or progress, until the plates were so corroded in spots, many of them of large area, that the hammer used in testing the plates broke through the skin of the boats under the effect of blows that would not drive a nail into a pure block.

The use of anti-corrosive or anti-toning paints, containing salts of any metal, is attended with the greatest charges to the coated structure. These pigments are extremely scientive to the presence of saline elements in moisture, their action being to tapidly dissolve portions of the iron, and to deposit the metal which they contain upon the surface of the plates, and these deposits exerting energetic galvanic action, cause corrosion and patting to go on with alarming rapidity.

Both merenry and copper salts are offenders in this way, but copper is by far the more objectionable, from the fact that the salts formed by the action of the sea water upon the compounds used in the compositions are far more soluble than the corresponding salts of merenry, and are therefore liable to be present in much larger quantity, and so exert comparatively a much more injurious action on the plates.

As an illustration of this, two equal portrons of sea water were saturated, the one with copper chloride, the other with mercurue chloride, and into each a piece of steel planed upon one side and of about equal weight and size, was placed and left tor locus days. At the end of this period the two plates were removed, and after being cleaned and dried, were again weighed, when it was bound that the one exposed to the copper-saturated seawater had left 2.2.2 per cent in weight, while the plate exposed to the mercurial solution had only lost 3.6 per cent, this being due to the much larger amount of the copper-salt soluble in the sea-water.

also suggesting various plans of attachment. In 1888 Mr. C. F. Henwood read a paper before the United Service Institute, strongly advocating zinc sheeting as attached by his system.

When the galvanic contact was small, then the sheeting had a certain life, but afforded but little protection to the iron, and gradually decayed away in a very uneven fashion; while in those cases where galvanic contact was successfully made, the ship on several occasions returned from her voyage minus a considerable portion of her sheeting.

Another drawback to the use of zinc sheathing is one which was found when it was used to coat wooden ships, and that is that sheet zinc, like every other metal, is by no means homogeneous, and that for this reason the action of the sea-water upo it, leaving out of consideration galvanic action, is very unevenly carried on, the sheeting showing a strong tendency to be eaten away in patches, while the metal itself undergoes some physical change and rapidly becomes britt e.

Attempts have been made to galvanize the iron before the building of the ship, but Mr. Mallett showed, in 1843, that this coating was useless when exposed to sea-water, as in from two to three months the whole of the zinc coating was converted into chloride and oxide; and when, therefore, galvanizing is used care must be taken to protect the thin coating of zinc. In any case the galvanizing must be done after the plates are riveted up, as any break in the surface would set up a rapid wasting away of the zinc, and the process could, therefore, be only used on small craft. Fresh water has less action upon the zinc than sea-water, and for this service galvanizing would be attended with some measure of success, the rapid wasting of the zinc in sea-water being due to the salts.

As has been before stated, if plates of iron or steel and one of copper be joined together or placed in communication and immersed in sea-water, acidulated solutions of water, or of mineral salts or oxides, the ferric body becomes electro-positive to the copper and is rapidly corroded. The corroded metal is always found in communication with the positive pole or current of electricity, the fluid soon becoming red from the rust formed.

tial and electro-positive, and none counteract cach certain but all attack the boiler metals.

The voltage in this case is distinctly recognized and evidently much different from the instances cited by Mr. Thomp on of metals under strain only. In one reported case, it was one obne, corrosion was marked and the current grew stronger as the corrector metalscal.

Fresh water and solutions other than was water, also vapors, are corrosive agents to boilers, the corrosion of which is modified, corrected and rendered uil, by the use of electrogens, or heavy cast zine plates. In old boilers using fresh or salt water, the corrosion in progress is arrested by the use of the electrogens, so long as any appreciable amount of zine is present. When the sine is wasted away or removed, "bleeding" from the boilers at once begins, particularly in old boilers or in the tube settings.

In marine boilers, sine 10" · 6" · 1" to the amount of one square foot of surface to 50 square feet of heating surface is placed in clean, firm, metallic contact with the internal steam or water surfaces. Too much sine is hardly possible and is better than too little. The amount of sine can be reduced after a time to 75 or 100 square feet. The sine must be placed in absolute contact with the bright metal at a number of points. Suspending the sine in any form in trays or baskets will not prove effective. Zane is slowly decolved in lot water, and deposited as a sediment that can be removed to the blow-off, carrying with it any old scale or rust loosened by the galvanic action of the zine. The boiler fluid contains a white flucculent precipitate of zine (zine oxide). If the water contains the subdistance or carbonates of lime or magnesia, silicates or other minerals that form the usual hard, vitreous scale, the precipitated sine exide unites with them and holds them in solution until blown out

Zine causing old boilers to bleed, might be considered an injury instead of a blessing. It indicates that the boiler needs repairs to prevent future disasters

The amount of zine in boilers for land service using waters containing mineral substances, has not been so clearly ascertained as in the case of marine work; but the results in the latter case are a good basis to reckon from,

of zinc and 19.656 parts of oxygen. The hydrogen set free replaces that lost in the heating of the water, that in a measure is broken up at all heats below a low red heat, where complete dissociation of the hydrogen occurs. In both the steam and water, the flocculent particles of the zinc readily unite with any ammoniacal, carbonic, or sulphuric acids, saccharine, or other organic vapors or liquids present to form sulphates, carbonates of zinc, etc., that would act mechanically in the water to prevent deposit and cause corrosion.

Central-heating-system pipes develop a virulent corrosion. The reason for some of the cases is difficult to find. This corrosion has

to the boilers, and in some cases the failure of the whole system. The corrosion occurs in both the steam-pipes and water-return pipes, being more marked in the latter. In screwed-end pipes the corrosion first attacks the heel of the pipe threads as zones of disintegration and extends until the whole pipe is affected, though no corrosion except at the joints may be noticed.

In the steam-pipes corrosion takes the form of pin-holing or pitting, from the inside at any part of the pipe and does not develop into a general corrosion of the surface as in the case of the hot-water return pipes. Whenever a bloomant or pin-hole from corrosion organs in

caused the abandonment of the pipes returning the condensed water

pipes. Whenever a blow-out or pin-hole from corrosion occurs in the steam pipe, a closing down of the metal around the hole by a peen hammer stops the leak, which seldom reopens. Peening the metal has made it more dense and ess iable to corrode.

Cast-iron steam-pipes are less affected than wrought-iron pipes, but the joints, draw bully, owing to the temperature changes that

but the joints draw budly, owing to the temperature changes that cause a leakage that frequent caulking only momentarily corrects. Cement joints are unreliable under high temperatures, while rust joints owing to their own corrosion burst the sockets of the pipes.

In central-heating systems, the losses from leakage and condensation amount to from 30 to 35 per cent yearly, aside from the loss of

the return water; while the corrosion losses are about 10 per cent of the cost of the pipe lines

Particles of dirt, einder in excess, unabsorbed carbon or manga-

[&]quot;Use of Zine in the Steamship Hindostan," Engineering, August 7, 1878.
"The Corrosion of Steamship Boilers." The Practical Engineer, Vol. X,

disintegration of the pipes, which continues after many of the above causes have been removed.

In all of these pipes a low density, open reasonst telement formed iron, einder in excess, heat, motion of vaped and water under pressure are all present, and no protective revenue of transmit. The hot water is more effective than steam in keepens the pipes clean and bright, ready for corresive influences.

The disturbance of water by high heat in being partially discociated has been already explained in this chapter. The action of zine on electrogens, in the case of matrix bories and sea-water corrosion, is always favorable for the preservation of the notal. The use of zine to prevent corrosion in steam paper, tachater and heating lines, could be hardly less favorable.

M. Londin's (Complex Rendus) experiments on the correspon of iron immersed in water usually found in feats booker, was. That with both ordinary and distilled water, the temperature had a very important influence, viz.:

"At 68° F, the quantities of oxygen absorbed par equate foot of iron surface per hour, when immersed in defalled water was 0.238 grain and in calcureous water 0.330 grain. At 24.2° I, the quantities rose to 2.364 and 2.579 grains. The immersion of non-in-water at pll ordinary temperatures was attended by the evolution of hydrogen, the action being the least in distilled water. At a temperature of 260° F, the decomposition of distilled water was equal to the alcorption of 0.01 grain per square foot of ferric surface per hour, and for calcureous water, 0.0129 grain. For water containing one fifth part of crystallized chloride of magnesium, corrosion was 0.0182 grain per square foot, and for water saturated with chloride of column, 0.05 grain; for sen-water of usual density, 0.067 grain, all per square foot of iron surface per hour immersion."

Corrosion Increased by Stress.

The tendency of iron to change its physical properties by a change in the condition under which it may be placed in ordinary attrictual work is strikingly shown by the following instance taken from Francisco April 27, 1921.

of the best foresinte from gave under test the following results:

"Tensile strength, 54,230 pounds per square inch; clongation on 8 inches, 28.4 per cent; contraction of area, 49.6 per cent. No traces of lamination were shown during the test, but some months after, the bar was found in the condition illustrated, which shows that it con-



Fig. 51.—Effect of strain on the corrosion of iron.

sists of layers of different chemical composition, those which have rusted being electro-positive to the other portions of the bar."

Iron rivets and iron plates in some cases show the rivets corroded and the plates unaffected, and sometimes the contrary, and so with steel rivets and steel plates; also iron rivets in steel plates or steel rivets in iron plates all show the most erratic evidences as regards corrosion, in many cases without reference to the character of the water used in the boiler or to the external conditions. As a rule, all analyses of the plates, rivets, and other material used in boiler work are made from samples as they come from the manufacturer's hands, and before being worked. Hence, when corrosion of either plate or rivet has attracted attention, it is seldom possible to get a sample of that particular make and lot of rivets to analyze to show at the the management of the the management of

caulking, etc. These processes, also punching instead of drilling the holes, develop corrosion, that takes the form of patting around the rivets and furrowing on the sheet joints.

Mr. Thomas Andrews, F.R.S., reporting to the Writish Institution of Civil Engineers, states his conclusions On the Enfect of Stress on the Corrosion of Metals.* In brief they are:

"That wrought iron and various steels, when exposed separately, without liability to galvanic action other than local, under the action of sea-water for long periods, showed a greater corro ion on the part of all the steels than the wrought iron; the advantage in favor of the iron compared with the steels amounting to 25 per cent and upward. It was also noticed that corresion was increased in the steels in proportion as the percentage of combined carbon was greater.

"It was found that the galvanic action between wrought iron and steels induced a largely increased correspon in both metals. It was also found that the upper and lower portions of a metal structure, or vessel, although composed throughout of the same metal, were exposed to electrolytic disintegration from the galvanic action set up by solutions of different salmity on the metal; conditions found almost constant in tidal streams, brought about by the gradual rise and inflow of salt water and the outward flow of fresh water; and there are strong evidences to show that magnetic inducate tends to increase the corrosion of metals

⁶ When, however, the strained metal is in galvanic circuit or combination with the unstrained metal in any solution, an increased total corrosion ensues from the galvanic action, which research has shown to arise consequent on the different potential between the two.

"It was demonstrated that stress of any kind considerably alters the physical properties of both iron and steel, by increasing their rigidity and rendering the metals harder, also greatly reducing their properties of elongation or ductility. It requires a higher tomage to break a strained than an unstrained bar of the same metal. A tensile stress applied to a wrought-iron shaft, that produces an elongation of only 2 per cent, increases the tensile resistance of the metal 2 66 per cent.

to metals aftered their structure, rendered them harder in nature, and more liable while in their strained condition to be acted upon by sea-water, or other waters, than in their ordinary normal or softer condition. The experiments, however, indicate that an increased total corrosion, in excess of the normal corrosibility of the metal, occurs in a metallic structure, from the action of the local galvanic currents which are shown to be induced between strained and unstrained portions of the same piece of iron or steel forging, bar, or plate. Hence a strain occurring in a metallic structure tends, owing to the local galvanic action thus set up, to increase any corrosive forces which may be deteriorating the metal of which it is composed."

The details of the experiments are: Pieces of iron and mild steel of known character were submitted to tension, torsion, and flexure strains, to ascertain the changes made in the metal, and if corrosive effects were in any number due to stress. For tension, a bar was strained in a testing machine until an elongation was produced of 23 per cent in three inches, and at the point of reduced area the bar was cut in two.

The halves were then turned down at the shackle or vise end, where they had been subjected to little or no stress, until they had an area equal to the end half at the point where contraction of area had occurred, both pieces being finished exactly alike and each piece represented a section of strained and unstrained metal. They were then placed at the same depth in a saturated solution of common salt to approximate the action of sea-water on metal, the immersed ends representing strained and unstrained metal. An electrical contact made between the two pieces of metal, through the medium of a delicate galvanometer (Thomson's), the difference in potential or corrosibility could be observed. It was found that in each case a sensible current was set up between the two halves of the specimen; the strained portion was in every case found to be the electro-positive element of the pair, corresponding to the zinc in a galvanic couple, indicating clearly that the strained metel was acted upon more rapidly by the solution, and more easily corroded than the unstrained metal.

The test made with specimens after being submitted to torsional

roded more rapidly by the sea water.

This conclusion was further supported by tests made with iron and steel plates, when a flat piece was compared with one bent into an U or semi-circular trough; the bent plate in each case proving to be the one most easily acted upon by the solution.

The experiments throw an interesting light on a subject which has hardly received the attention it deserves, and helps to explain some of the peculiarities in connection with the wasting of certain structures that have been involved in considerable invitery. The metals operated upon by Mr. Andrews were large, redied wroughtiron bars and hammered wrought from shart. Bessenier steel and Siemens steel forged shafts, also arge into of soft and hard Bessenier and Siemens steel; soft and hard east steel, and steels made from each of the metals aluminum, nickel, silicon, and copper. Experiments were also made on rolled plates of wrought from soft Bessenier and soft and hard Siemens steel and soft east iron. The chemical compositions and general physical properties, etc., of all the metals are given and talanlated. All the metals experimented upon were perfectly bright.

General results: The average electromotive force obtained by-tween strained and unstrained portions of the same metal were, viz.:

Wroughtsiron forged shutts	() tile.	seelin.
Soft Besseiner steel terged.	11 11111	**
Hard " " "	(1 1 ta 14 s	**
Soft cust steel	## #### <u> </u>	èe
Hard " "	11 1813	**
Silient steel	** ****	**
Aluminum steel	11 11111	4.6
Nickel steel,	18 CH 8.1	**
Rolled wrought-iron bars,	11 1817	**
Soft Siemens steel	11 (11)	**
Hard " "	11 1815	**
Copper steel		**
Chrondun steel,	() (H1)	11
Bessemer steel lummered forgings,	. n uii	it
Siemens steel " "	ii inni	**

With cold-drawn small steel rods in galvanic circuit with copper

than in the tensile, also in the flexure, tests.

These electric measurements ought, perhaps, to be regarded as tentative indications, establishing a general principle, rather than as an absolute measurement for the purpose of accurate comparison of the behavior of the various metals. The chemical analysis of all the metals was made prior to straining them. These experiments extended from a few seconds to over ten days, in which it was observed that the difference in the electromotive force between strained and unstrained metal steadily declined from the initial amount, but was in no case extinguished.

Corrosion, or the oxidation of substances by chemical action is always accompanied by electrical energy, that may be of more or less intensity, or electromotive force according to the substance consumed.

Chemical action is probably due to the unbalanced attraction among the various molecules of matter lying in juxtaposition, the rearrangement of which caused by strain or a change in the thermal or electrical conditions of one atom changes them all. It is known that a change in either the thermal or electrical conditions develops corrosion in certain circumstances but does not in many other cases of apparently the same nature. The amount of electromotive force developed in the oxidation of a few substances is indicated in the folowing instances:*

	is a red congruence that is a second	and a street contribution of the contribution	
Sulatnines.	Heat of Oxidation of Equivalent, Calories, B. T. Unite	E.M.F. Relative to Oxygen.	E.M.F. Relative to Zinc.
	••	, wa-me-	1 th following rate in 11 11 to report a content displacement of
Carbon,	2.000 - 7.938	(1, 09)	1.7.1
Silver	9,000 - 31,742	0.39	-1.44
Copper.	18,760 71,057	0.80	1,08
Land	25,110 - 99,016	1.12	-0.71
Iron,	31,120 135,415	1.55	-0.28
Zine.	12,700 - 109,071	1.83	0,00
Peroxide of lend -	12,500 - 48,022	0.52	-2.35

The corrosion of ferric bodies results from the decomposition of water or air by electrical energy.

As detailed in Chapter III, atmospheric moisture in the presence

black magnetic or stable oxide of iron, that in manufactured articles is represented by the Bower-Barff products.

Every pound of iron oxide represents the energy of 1.668 pound of coal required for its formation. This rust requires .3375 pound of water to furnish the necessary oxygen. A pound of iron oxide represents the corrosion of 13.43 square test of metallic iron $_{1000}$ inch in thickness and has developed 163,233 B. T. Units (.0641 H.P.),

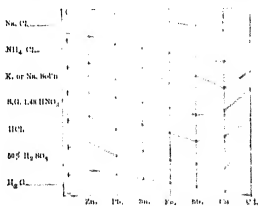


Fig. 52. Diagram of the corresibility of metals,

or equal to an electromotive force of 1.55, to neutralize which would require the oxidation of .847 pound of zinc.

In all of these exidations the thermal manifestations are subordinate, and, with the electrical energy being of low potential, when they extend over any considerable period, are unrecognized or difficult to determine.

The affinity of oxygen for hydrogen represents an electromotive force of 1.47 volt. The decomposition of water acidulated by sulphuric acid yields at the cathode 11.12 parts of hydrogen and at the anode 88.18 parts of oxygen; about 1 per cent of ozone being formed from the oxygen.

The decomposition of 1 pound of zine for the protection from corrosion of murine boilers or other like ferric hodies evolves 9840 cubic

		B. T. Units.	Horse-power.
Bel 1	According to the second second second to the second		
		900	0.35
	of copper) battery,	1419	0.55
Grove's or Bunsen's	(nitric acid)	2722.4	1.06
Peggendorf's (chrom	ic ncid)	2827.5	1.14
Solohorie acid (1 in !	parts of water)	3006	1.17
			1

All losses from resistance being excluded,

A series of experiments covering several years, upon the corrosibility of metals, has been made at the University of Wisconsin, under the supervision of Prof. Dugald C. Jackson.

Prof. Jackson, in the discussion of paper No. 901, "Protection of Ferric Structures," * referred to the results of his experiments, from which I briefly quote: "When a piece of iron or steel is placed in a testing machine and its electrical condition is followed up during the straining test, its corrosibility appears to increase practically in proportion with the strain, so that a diagram plotted with stress along one coordinate and corrosibility along the other appears to be of almost exactly similar character to a diagram plotted with stress and strain along the two axes of coordinates.

"Two illustrations of these diagrams are presented, Figs. 53 and 54, from test pieces of wrought iron

"In the case of cast iron Fig. 55 shows the stress-corrosibility diagrams for two specimens in tension. A comparison of these diagrams with those for wrought iron in tension, illustrated in Figs. 53 and 54, shows the marked difference between the two metals. Fig. 56 shows a stress-corrosibility diagram for cast iron in compression. The exact forms of the diagrams taken from cast iron depend in some degree on the physical character of the specimens, but the diagrams shown are typical ones. The effect of strain is small in the case of cast iron.

"The corrosibility of the specimens was measured by determining the electromotive forces of the test pieces toward a standard electrode in a normal solution.

"The results of the tests show that in bridge members and similar pieces that have been worked, the metal appears to be easily affected by corrosion, this corrosion being properly characterized as

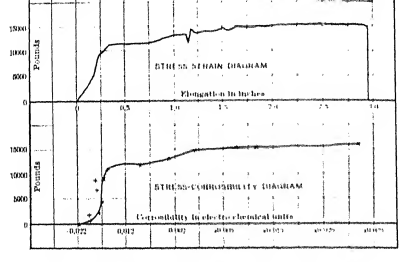
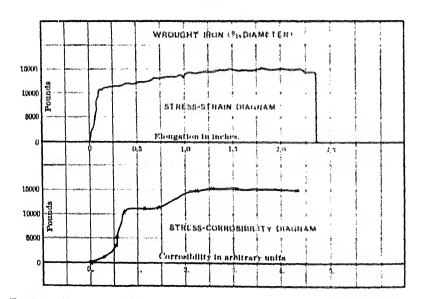
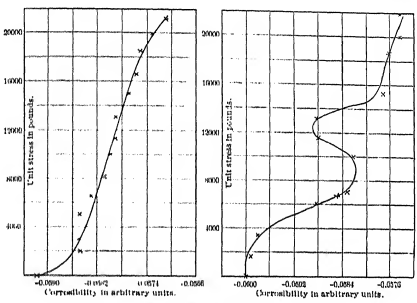


Fig. 53.—Stress strain and stress corresibility of a rought iron.



away and the unstrained metal is not.

The experiments give a satisfactory explanation of much of the so-called grooving in boilers and corrosion of a similar character. Here the strained metal of a punched boiler plate that is not completely covered by a rivet-head becomes eaten away. Or perhaps a plate becomes strained at a joint by temperature stresses and the strained



Pro. 55,-Stress-corrosibility diagrams for east iron.

streak is corroded. In each case the strained metal is of greater corrosibility, and it acts as one of the plates of an electric battery in which the other plate of the battery is the unstrained metal of the boiler shell, and the electrolyte is the water within the boiler. The strained metal is the electrode which corresponds to the zinc of the ordinary voltaic cell, and it is eaten away.

"Another illustration of corrosion of this character is the so-called (by bridge engineers) Cooper's lines, which are often evidenced in the corrosion of bridge members. These are lines of electrolytic the state of the s

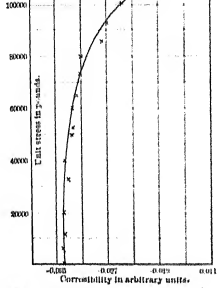
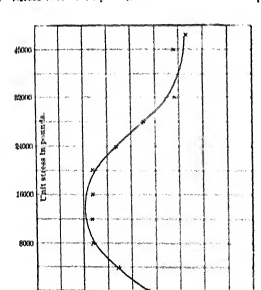


Fig. 56.—Stress-corresibility diagram of cast non-in-compression.



"It is also true that metals which do not change their physical characteristics when strained, apparently do not materially change in corrosibility as the result of strain. Thus if lead is stretched, its corrosibility does not appreciably increase. The same is true to a certain degree of bruss and copper. Fig. 54 represents a stress-corrosibility diagram for hard-drawn copper. The same is true also, to a limited degree, of very soft iron, but as even the softest iron does harden sowewhat when strained, its corrosibility is somewhat

An experimental study of the corrosion of irou and steel under different conditions has been made by Mr. Carl Humbuchen, B.Se.* These experiments were conducted on the lines of those of Mr. Thomas Andrews, F.R.S., hereinbefore given, "on the effect of strain on the corrosibility of metal."

affected by strain."

Hambuchen's apparatus and method of testing, and the checks and precautious against errors, especially in obtaining the value of the electromotive force, were superior to those heretofore employed by experimenters, and the results are more in accordance with the practical experience of the present day.

A hindrance to experimental investigation of the corrosion of metal is the length of time required to produce measurable results. Hambuchen took advantage of the fact that corrosion being developed by an electric current flowing from the ferric body to the electrolyte, the rate of corrosion could be greatly increased by causing a current generated externally to flow from the metal as an anode, thus causing the corrosion to occur under what may be termed exaggerated or intense conditions, the metal being corroded as much in a few hours as it would be in as many years by exposure to the weather; the resultants being practically the same as the effects produced by ordinary corrosion.

The losses in weight from corrosion in different irons, steels, and other metals, under strain, from nil to breakage, are tabulated, also the loss in weight of metal per ampere hour, and the electromotive force developed at various points of the strain.

Normal solutions of ammonium chloride, ammonium sulphate,

the metal in the different solutions was practically the same, and that whether the salts were sulphates, intrates, or chlorides shed not materially affect the rate of corrosion. The ampete four stand from 11 to 13.6 and the exposures from 19.5 to 24.25 hours.

Interesting facts were developed showing the variable inture of "pitting" in mild steel exposed to different solutions. Thes, 58 and 61 show a round pitting as the result of the anim onium chloride; an clarificated pitting results from the ammonium sulphate, and a more uniform corrosion from a potassium intrate solution. In the cust-iron specimens, Fig. 59, the corrosion consisted of a soft carbona.

olocharace, the experiments determined one on a set in result of

an elangated pitting results from the animonium surphate, and a more uniform corrosion from a potassium intrate solution. In the cast-iron specimens, Fig. 59, the corresion consisted of a soft carbonal econs uniterial, which generally adhered very fittidy to the surface of the iron. In case the current density was carried beyond 10.025 amperes per square inch, this soft material would separate from the iron after attaining a certain thickness. The formation of this layer must offer some resistance to the flow of the current, and therefore protects it to a certain degree. Or, in other words, a given potential difference between an iron pipe and a railway track would cause less flow if the pipe thus conted were east iron, than it it were wrought iron, which would quickly reveal its weakness by the difference character of the corrolled conting. But with a given uncount of current flowing from normal cast iron and wrought iron, the corresion is nearly equal in amount in the two.

It was noticed in the case of the cast iron anodes that there was

It was noticed in the case of the cast iron anodes that there was a liberation of gas during the process of corresion. That this was not due to the flow of the current was shown by interrupting the current, the liberation of the gas continuing for some time after. The nature of the gas was not determined, nor of what the action consisted.

If the current density was not excessive, the iron did not undergo any material change in appearance, even though subjected to the action of the current for a long time. But although the general form and outward appearance of the east iron remained the same, the fact that its structure had been materially altered was shown by cutting it. The east iron was found to be softened to a certain depth, the material removed having the appearance of face iron plates of the same metal.

Changing the crystalline structure of steel by annealing, hardening, and burning, causes the amount of corrosion to vary, as is noted in the tables on pages 363-364. The amount of corrosion per ampere home of the hardened steel is considerably less than that of the

annealed or burned steel.

This is an apparent discrepancy with other data and observations in regard to strained metal being the most subject to corrosion. Hardened steel, being necessarily under a highly strained condition, should have shown greater corrosibility than the annealed or burned specimens. That it did not is owing to the fact that the high tension between the particles held them in place until they were severally corroded entirely away. In the burned or annealed steel the particles, when only partially corroded, were loosened in their bond to each other, and east from the mass before being entirely corroded. The pitting of the annealed steel, Fig. 66, the composition of which is similar to that of sheet iron, Fig. 72, shows a greater corrosibility than the hardened steel, due to the above reason.

The small percentage of carbon and other impurities in the steel would not account for the corrugations in the burned and hardened steel, shown by the figures. It would be unreasonable to suppose that the impurities or curbon could be regularly distributed as indicated by the corrugations; they must be due to lines of strain in the metal that corrosion developed.

The metals that are electro-positive to iron and steel are magnesium, aluminum, zine and cadmium; while lead, antimony, tin, copper, silver, carbon, manganese, and some of the metallic oxides, are electro-negative to ferric bodies.

Test plates of clean, beight wrought iron, cast iron, and steel were drilled and the several holes plugged with one of the above metals. The plates were their placed in sand saturated with animonium eldoride and other corrosive solutions, and after a short exposure were examined; all the plates in which the electro-positive metal plugs were placed were found clean and bright, while the plugs were more or less corroded. In the other set of plates the surfaces were

SPUC68" (UC GILLOUCHO ROZER PLANT THE DIFFER TO THE SOUTHOUT WHAT (UC plates were corroded. The conditions of electrolytic corrosion apply to most of the metallic oxides as well as to the metals, and are developed in both;

1. When two or more conducting substances are in contact with an electrolyte, 2. Whenever there is any difference of electrical potential between such bodies. 3. When a suitable connection between the conducting substances furnishes a path for the flow of the current.

All of these conditions are present in the decay of paint contings. as well as the corrosion of iron and steel. The electrolyte consists of moisture in any form, and may be acidulated, spline, or field

Iron and steel rea never pure or homogeneous, they contain upon their surfaces many substances, such as carbon, graphite, millscale, and particles of metal and oxides. The body of the metal may be formed from scrap iron of different natures, and the heat of fusion seldom renders the mass homogeneous.

In cast iron and steel, in their many processes of manufacture, there are many irregular zones of density and purity each of which has its own potential. Between all of these differently charged bashes a current of electricity is set up, the circuit being completed through the electrolyte.

This electric current flowing from the metal to the electrolyte will cause corrosion of the metal, which may be general over the whole surface of it, or be localized in spots, according to its composis tion, which is affected by local disturbances, such as welds, strains, annealing, burning, and hardening, or the presence of foreign safestances. Some peculiar cases of corresion can be explained by ascertaining if the metal has been subjected to some of these influences, that otherwise would be classed as mysterious.

Mr. Hambuchen, in order to draw a comparison between electrolytic corrosion and ordinary corrosion, immersed specimens similar to those exposed to electrolytic action in a tank containing a normal solution of ammonium chloride, and left them undisturbed for four months. The results obtained showed that the amount and character of corrosion depend upon the quality of the metal, and confirmed the conclusion derived from electrolytic corrosion. The time of exposure of these specimens was, however, too short to develop any marked pittings an other assumits affects the second of the second

actal which are similar to those produced by corrosion under ordinary onditions.

In many if not all cases the character and rapidity of ordinary orrosion of iron and steel depend upon their physical and chemical respectives, and the galvanic action due to differences in potential between different parts of the metal.

The application of stress to metals causes an increase in chemical activity, this increase being especially marked after the elastic limit s renched.

It is possible to plot a curve showing the relation of electromotive orce to strain, which is similar to that of stress to strain.

There is a definite relation between the electrical potential of my metal toward an electrolyte and the amount of energy stored up at the metal through the application of stress. It is evident that the protection of ferric structures from corrosion requires their removal from electrolytic influences.

The several sperimens subjected to different conditions of corrosion were all taken from the same bar or sheet to facilitate comparison.

The following tables are means of a few of the separate results given by Mr. Hambuchen:

Table showing the less in weight of iron and steel used as anodes, immersed in a solution of announum chloride and exposed to the action of an electric current of varying densities and time.

Comment of the Commen		1 1		}	
Material and Condition of Surface.	Area in Equare Inches.	Total Loss in Weight. Grams.	Weight Lost per Ampere Hour, Grams.	Ampere Hours.	Ex- posure Hours.
Annealed steel, polished	10. 10.523 10. 10.48 10.3 10.555 10.073 10.093 10.373 10.203	15.9 15.33 14.633 14.407 15.666 15.515 12.875 12.273 9.996 13.87 14.63	1.1883 1.1163 1.077 1.059 1.1526 1.1575 1.1475 1.0983 0.7506 1.184 1.084	13.6 13.3 13.6 13.3 13.6 13.3 11.2 11.2 13.3 11.2 13.3	23 \$\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\
		Tonger .		•	

* *	THE REAL PROPERTY.	* 1

Motal and Condition.	Apple see Properties	Lorent Loon the Wingston Colonia	Wraght per square linh.
Sheet iron, surface polished. "" scale partly removed. "" with scale Cast iron, surface polished. "" scale partly removed. "" with scale. Annealed steel, surface polished. "" scale partly removed. "" with scale. Hardened steel, with scale. Steel with scale, burned not hardened.	10 1.17 10 30 10 556 13 506 14 16 14 117 10 42 10 603 10 37 10 57 10 255	11 (7.14) 1	0 066 0 142 4 698 0 0793 0 117 0 1174 0 0708 9 102 0 193 0 198 0 3613

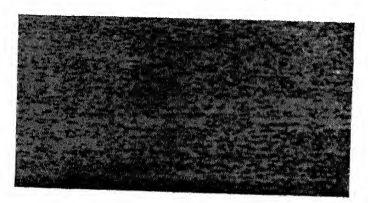


Fig. 58, - Mild steel (ammonium chloride solution) — Magrate d 23 dramaters,





Fro. 60. Cast iron (with scale). Magnified 2½ diameters.

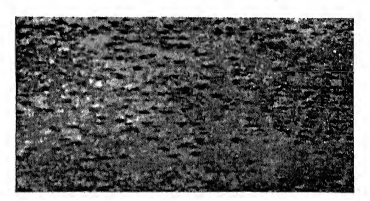
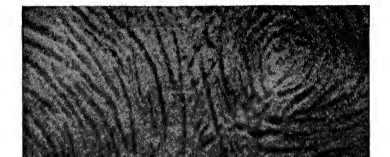


Fig. 61. Mild steel (ammonium chloride solution). Magnified 24 diameters.



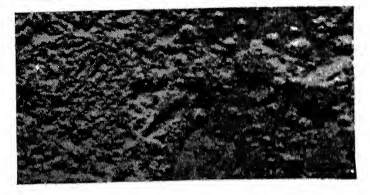


Fig. 63, --Burned steel not hardened (with scale). Magnified 24 diameters.

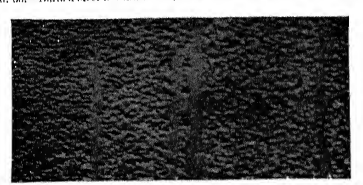
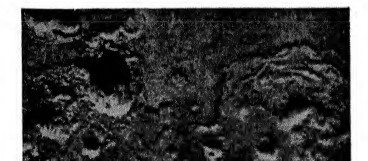


Fig. 64.—Annealed steel (polished). Magnified 24 dismeters,



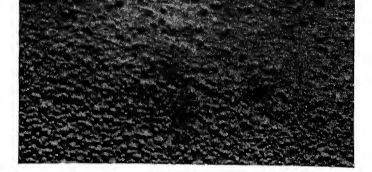


Fig. 66.—Steel burned but not hardened (polished). Magnified $2\frac{1}{8}$ diameters.

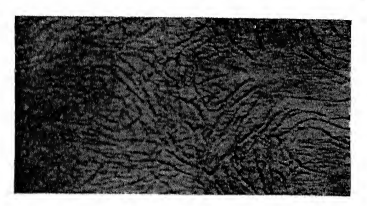
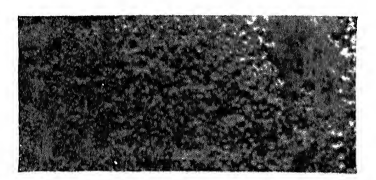


Fig. 67. Hardened steel (polished). Magnified $2\frac{1}{8}$ diameters





Fig. 69.—Steel burned and hardened (polished) - Magnitud 21 diameters.



Fro. 70.—Sheet iron (polished). Magnified 21 diamsters.



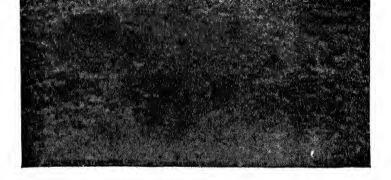


Fig. 72.—Sheet iron (with scale).—Magnified $2\frac{1}{8}$ diameters

CHAPTER XXXIV.

PERCENCIASIS OF UNDERGROUND MUCCUS.

Proc. Edwin J. Houston defines electrolysis as chemical decomposition effected by means of an electric current; that is, the source of electrolytic corrosion, lies in the release of an atom of oxygen from any moisture and its instantaneous combination with any iron it can seize upon."

Prof. D. C. Jackson (University of Wisconsin) defines its action:
"In an electrolytic cell with iron electrodes, having any salt or salts of alkalino metals or earths in solution in the electrodyte, the salts are decomposed, their acid radicals attacking the anode, for one an iron salt.

A proof of this theory is found in the storage battery, in the charging of which a large current is discharged from a lead plate into an electrolyte consisting of dilute sulphure acid. In a storage battery the exidation of both plates takes place, but the red exists of lead formed is very different in character from the corresive effects of an electric current on an underground lead pipe."

*"There is nothing mysterious about the corrosion of metals by electrical currents. Its action is precisely similar to that employed by electrosplaters in their art. Two plates of metal placed in any material, whether damp earth or the solution vat of an electroplating apparatus, a current of electrical energy will be instituted from one plate to another and the plate or object from which the current flows will be corrocled. The current will make its own selection as to its course and the body to be attacked. In the case of metals buried in earth or water, the electrolytic action is out of sight and probably out of taind, but none the less present and uncontrollable; while in the plating bath it is in sight and controllable more or less, at the will of the attendant. In the case of high voltage and large ampere currents returning to their source of generation, it is

current will divide and return to its source in the direct ratio of their conducting capacity, whatever these conductors may be. Even with large and well-bonded metal return conductors buried in the earth or in conduits, some of the current will invariably pass by way of the earth and reach any outlying metallic bodies. The low voltage of .001 to .01 and the amount of amperes will determine the rate of corrosive effect in all metal in their course."

The advent of the steel building almost simultaneous with the introduction of the dynamo, has added not another form of corrosion, but a new field for its development and a new danger.

The principal part of the metal in steel frame structures is so embedded in masonry as to render inspection of its condition almost impossible. The pipe systems are more accessible, but are nevertheless at all times a ready prey for electrolysis.

Three hundred horse-power of electrical energy are not uncommon installations for light and power in one building. Whether led in from the outside, or generated in the building does not change the effect, which is to disturb the normal electrical conditions of all metals in the immediate neighborhood and in many cases those far distant.

In the return of this energy from its work to its generating source, if the pathway is not unde perfectly free by the use of a conductor of adequate size, or if it be of such a length as to render a shorter and better circuit through other objects possible, then the current will jump the line wholly or in part. On the new route, wherever it leaves the metal, another jump will occur, and the metal will be corroded at that place, and not where the energy entered. There is always moisture enough in any building to afford adequate oxygen for the corrosion.

In the steel frame work of the building, the electrolysis will be at the foot of the columns nearest to the least resisting pathway of the current, generally at a point impossible to locate or inspect, and with moisture in excess to make a good cross-cut and dangerous circuit.

In the iron pipes there usually will be a jump at every joint, if it is made by the hemp gasket and lead, or by a hydraulic cement method. have been found to be the cause of a jump corresion, particularly if aided by the presence of damp dirt or other substances.

A voltage of 118 maintains an arc of one inch in an electric arc light, and requires 1 H.P. of electric energy, and fourteen sixtoon-candle-power incandescent lamps represent the same amount of energy. It is not infrequent to find an arc light or twenty or more incandescent lights, or an equivalent energy from small motors connected to the pipe systems or the steel frame of the building. The effect of these strong currents is to set up a corrosion in the steel at some point where the current is interrupted on its return to the dynamo.

Under some one of their many developments, induced currents are strong enough to correde metal, even if the main current or return current wires are adequate for their duty. Less than .005 volt establishes electrolysis, the amount of which is in proportion to the amperes present and not to the voltage.

Protection from the effects of the jump of the current, particularly in the lower parts of a steel frame, is rendered more uncertain, by reason of the disturbing action of electric currents from adjoining sources of generation. These currents finding then return to their source of generation obstructed from any cause, form a short or easier circuit that often lies through another dynamic's sphere of action. They invade its field and disarrange a return current system that at first might have been adequate for its duty, but is not able to withstand currents from its neighbor of subsequent installation

Twenty or more of these electrical installations of shifterest degrees of voltage, ampere, and work, are often placed within a comparatively small area. Many of these have been found to have a return wire system of inadequate size or of faulty insulation, and all of them subject to a wide range of fluctuation in energy due to the varying character of their separate work. With these influences at work at nearly all hours, it may be confidently expected that the near future will reveal some large and dangerous examples of vorticeous in steel

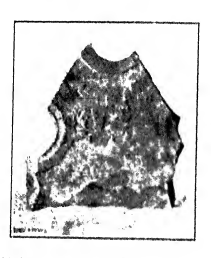
some three-fourths of a mile distant. This branch of science so closely associated with the daily needs and welfare of mankind, was paralyzed by the culpable indifference of a corporation to the requirements of science. It took an Act of Congress, carrying the imposition of a heavy fine to stop the nuisance. The Act not only prohibited the use of any underground water- or gas-pipes as means for returning the trolley-line currents to the power house, but also, forbade the connection of either pole of a railway dynamo in any direct manner with the earth.

Paints furnish neither renedy nor protection from electrolysis, Paints under catchy na ness are extensively advertised as being electrically inert, or insulating in chareter. Such names and statements are misleading and unreliable. No paint whose pigment is an oxide and again reducible by heat to a metal, is non-electric or passive to electrical influences in any degree beyond that due to the difference between the oxide and its metal, generally about 50 per cent, but is never uit.

Lampblack and graphitic carbon are the only pigments that are partially non-electric, and even with the use of these in a paint, the coating as an insulating substance is governed by the vehicle. The vehicles containing the resins, fossil gams and refined bitumen and combined by heat into a variash, are the best for non-electric paints. It is quite amisual to find them in use on account of their cost, while the cheaper grades of resins and resin-oils used in the vehicle are only insulating up to a certain percentage, when they become conductors.

All of the vitreous class of pigments, such as slags, hard-burnt brick, tiles and slate, are conductors in their pulverized form, and usually act in a paint as the negative electrode to concentrate the electrical energy upon the covered ferric body. The thin coating of the vehicle, $\frac{1}{4} \frac{1}{a_{nn}} \frac{1}{4} \frac{1}{a_{nn}} \frac{1$

Electrolysis inaugurated beneath such coatings, generally throws them off, or they act as a mask to conceal its ravages; while inferior 1900, with the loss of life of two persons and the injury of fourteen others. The stand-pipe was 60 feet away from the other stand-pipes, and more than a mile away from the power station. I ig. 73 represents a sample of a steel sheet from the stand pipe, showing the pitting in the sheet around the edges of the rivet heads. The examina-



Pio, 73. Pitting of steel statul pape short around the risets

tion of the wreck of the stand pipe showed that the whole usner surface of the vertical shell appeared to be thickly exerted with blisters resembling in outward appearance the tubercules sometimes found inside of old east-iron mains.

A similar stand-pipe on the East Bluff was drained and was found to be similarly pitted. This blistered covering, which was almost as thin as paper, was composed entirely of covide of tron, and on brushing it away the black paint with which the stand-pipe had been originally coated was found beneath it. The paint was oftentimes almost unbroken, or, at least, very slightly cracked. When the

^{*} Excepts from "Electrolysis of Underground Metallic Structures" A paper rend by Mr. Darney H. Maury, Chief Engineer of the Process III Water Works, before the American Water Works Assessment Mrs. Lines & A. Lines & Co.

paint was brushed off the pit would be disclosed, considerably smaller in area than the surface covered by the blister. The surface of the metal in the pit was perfectly bright and clean and its fibre was clearly discernible. Many of these pits were more than 1 inch in depth. They were slightly more numerous in the West Bluff standpipe than in the East Bluff standpipe, and were in both generally larger and deeper on the lower courses of the vertical shell.

The electrical examination relative to the stand-pipes was conducted mainly at the East Bluff stand-pipe, which was still in service. A flow of a part of the current from the railway-line was clearly traced through the earth to the anchor-bolts which held the stand-pipe to its foundation, as shown in Fig. 74, up these bolts and into the steel of the shell, and through the shell and from its inner surface to the projecting section of the 16 inch flanged east-iron pipe which served as both inlet and outlet, and which connected the stand-pipe to the water-mains. The current was then traced along this pipe and along the nains to the power station. The deflections of the volt-metre needle were clearly traced to the railway current, being especially influenced by the cars on the line beyond the stand-pipe. and when the cars stopped running at night, the movement of the needle ceased. Where the current left the inner surface of the shell to pass through the water to the inlet pipe it made the pits already described.

Fig. 78 shows the interior surfaces of three sections of this inlet pipe, marked A, B, and C, respectively, the positions occupied by these sections originally being shown by the letters A, B, and C in Fig. 74. An examination showed that strongly marked and numerous pits were inside the sections A and B, while the inner surface of the section C was practically as smooth and perfect as though new. When the condition of the inside of these three sections of pipe was first noted, it was hard to understand why A and B should be pitted, while C was unaffected. A closer examination, however, showed that in the flanged joints between the bottom sheet of the stand-pipe and A and B, respectively, corrugated copper gaskets were used,

pipe and water-mains to the power station, encountered, at the join

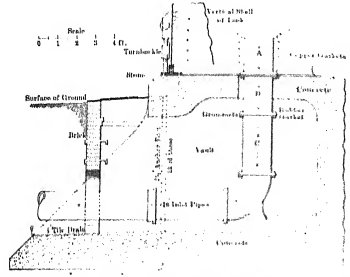


Fig. 74.—Partial section of Peoria stand pipe, showing course of electrical turrent.

between B and C, the rubber gasket and the granuacts. The effect of the gasket and grunnets was to practically in plate the section C



Fro. 75. Interior view of lengths of sidet to stand paper showing pattings in A and R, and effect of the simulation of the

oxide on the inside of the pipes around the joint between B and C, in order to continue on its journey. As the current was not leaving C, this pipe was not injured, but the current, in leaving the inner surfaces of A and B, did pit them, as shown in the photograph.

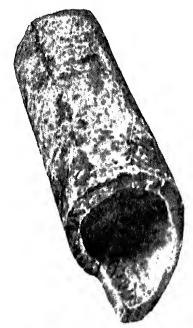
The experiments conducted since the destruction of this stand-pipe have determined that no manner of packing the joints in an underground east iron water or gas-pipe line affects, only in a small degree, the difference in potential between the two ends of the connected pipes. Whether the joints were well or poorly calked, the pipes empty and dry, or full of water, clear or muddy, with scale on the pipe or clean surfaced, the drop of potential around the joint only varied from 0.0145 to 0.008 volt, and the general average resistance of the joints was about 96 per cent of the resistance of the whole line of pipe, and the resistance of the joints increased with age. Pitting was always observed where the shunt of the electric current



Fig. 76. Electrody to all pattings on 12 inche east from water-main. (Current flowing from A to $B_{\rm c}$)

left the metal to flow around the joint, and this corrosive action was as marked upon the inside surfaces of the pipes as upon the external surfaces; but from the conditions could not be observed. Wrought from pipes joined by the usual screw-thread and thimble connections were almost as universally attacked by electrolytic action at the joints of the contains a times. The difference in retential

which generally showed in the form of blisters or like the usual tuber-



Fro. 77. Electrolytical pittings no monde and entends of 4 meli cast iron water page

cules formed on water-pipes by the action of water. There tubercules when broken, and the material under their analyzed showed 22.3 to 23 per cent graphitic earlien and 47 to 47.7 per resistances.

The soil in which the pipes were embedded had received and was impregnated with the metal tiron or leady corroded by the electric errent, even to some distance from the pape. In sections of pipe from 600 to 2000 feet long, there were reveral rotan and centres of greater action than on the pipe in general but there all centred toward the nearest generating dynamic, and were explicitly localized at these points, from the greater amount of electrical energy east off at the meeting and passing points and switching of the ears, contiguous lines of ears and currents from other sources all contributing in the most erratic manner to the corresive result.

service pipes for water and gas, laid in a light, sandy soil. The water service pipes corroded in seven years, so as to need renewal, and the gas service had almost completely disappeared at the end of ten years.

Analysis of the soil disclosed the presence of common salt, magnesium chloride, iron, alumina, silica, lime, and posphates. The town where the pipes were laid had no system of sewers, and during the summer senson the streets were daily watered, and as the streets were level the water was quickly absorbed. The subsoil in which the pipes were laid was persons and alternated from dry to damp.

Carbonic acid was generated from the chemical action of the soil and attacked the pipes as stated. The trouble was confined to the wrought iron service pipes; the east-iron pipe-mains to which the service pipes were connected were not affected in any noticeable degree. Similar pipe services laid in neighboring towns where the soil was of a decomposed granite nature were comparatively uninjured after a period of twenty, even years.

Salt or line in any soil in which pipes are laid necessarily prove active agents to promote corrosion, as they are hygroscopic in nature; and if alkaline substances are also present in the form of ashes and coal cinders as they nearly always are in the soil of towns, the life of all wrought non-work buried in it will be very short, even with the name crude coal tar contings. Clay puddle around such pipes proves a good protection, as pipes so protected have been found tractically uncorroled after forty years.

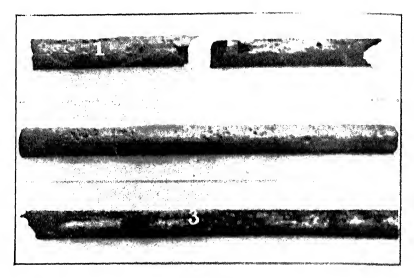
Mr Chas W Rowe, Secretary of the Dayton, Ohio, Water Department, reports that the east iron water-mains laid in 1891 were found in 1909 so greatly affected by electrolysis as to endauger the water supply of the whole city. Voltages of 4.5 were found in many parts of the paper system.

The Annual Report of the Water Department shows that in 1899 579 feet of 6 meli pipe and 26 feet of 4 inch pipe were abandoned on account of electrodysis.

Mr. Rowe reports that in 1898 over 46,000 feet of the waterpipes from 4 to 16 inches in diameter were so seriously corroded by electrolysis from the trolles line currents, that in some cases over the electric current used to drive the trolley cars was found to pass through the pipes and only \mathfrak{gl}_0 passed by the street rade on its return to the dynamo.

Fig. 78 shows the electrolysis of a 4-inch cast iron water pipe at

Reading, Pa.



Fitt. 78.

No. 1. Laid 22 years or about 18 years before the advent of the trolley lines at the point where the pipe was last

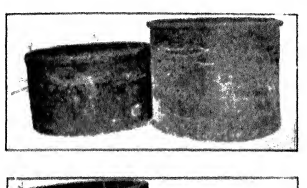
No. 2, Laid thirteen months.

No. 3. Inid 22 years. Burst when muonered for examination.

Fig. 79 shows a 16-inch cost iron suction pipe, also a water main from Rending, Pa. The pipes are samples of those Last 100 son and 1000 feet distant from the street troller lines, and all some over two miles distant from the electric power station. The effect of the resistance of the joint packing is seen in the estreeties of the ring on the end of the pipes; also in the general correspond at the ends where inclosed in the hell

Fence unils were driven into the ends as easily as into a securior

Kelsey, Chief Linguiseer, of Sult Lake Cits, Utah, is a condensed report of the reports from the chief engineers of seven eitnes in the United States, of the presence of electrolysis in their water supply systems,





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Briefly offsed relectival currents of hew potential, fill volt, were found and were enough to contain the electrolysis that under favorable conditions for the pipe might not have become serious in a limited period, but in the case of mile carrying units and and neith of decomposing materials the correspon from electrolysis was naterially necelerated and increased rappills no the voltage two to the volt. Currents of one half yell because of a telephone make in a few months. A minute quantity of solide outs in the outside outside outs the current and establish the electrolysis points in the metals, the correspond of which continued as long as the current flowed. In all cases where the mater pipes or the netal work of the building was positive to the earth or any corresponding object, the electric current flowed in that direction and electrologics was established in the period

ive action upon lead is of interest, as metallic lead has been ally considered electro passive, and is used for the outer insulativering for the cables in all underground conducts for electric g and power.

e Report of the Board of Commissioners of Electrical Subways,

st the pressure of the water when it was over 20 pounds. This

e Report of the Board of Commissioners of Electrical Subways, lyn, N. V., 1894, states that nearly 300 miles of lead coated one cables were rendered useless by electrolysis from the trolley is in that year. Many cases of correspondence to troin where all cable was incased in pitch and other insulating compounds.

A. A. Kandson, E.E., reporting the electrolysis of a 48 meh for water-min in the city of Candondge, Mass. Found volsof 25, and uniperages of 30–50 strand 90 at many points of the supply system.

R. I. The pipe was brinch thick when last and had been in

seven years.

10, 80. Electrolysis of a filiarle rant from page at Providence II I

. SI shows the electrodysis in one end of a read transfor a railway bridge at Providence II I. Hart made of the stranger

under the law of divided currents, would invariably leave the rails and seek another source of return through near-by metal. The smallest measure of difference in potential between two metallic bodies was sufficient to produce and maintain electrolysis in one of them."

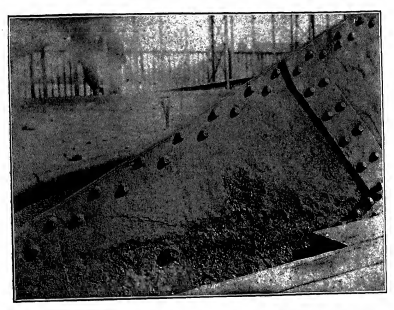


Fig. 81.—Electrolysis of a steel bridge truss.

Dr. Leybold's paper, "Electrolysis of Gas-pipes," * states: "The pipes when laid were protected with canvas soaked in boiled coalgas tar, and the destruction of the pipes was more rapid than where they were laid without the canvas coating. New pipes laid with canvas and tar coatings to replace the old ones were perforated into holes in seven to eight months."

The annual report of Mr. Wm. Jackson, City Engineer of Boston,

More calls attention to the correction of 48 more mains for water mains in that city, that have been cated away in mains phase but one half inch, and could be cut as easily a physician. The diametrations from the bursting of such a paper inapparent.

The practical effect in the corresion of an iconomial gas mains and wrought from service paper, principally the color mail stranger, is noted in the Official Reports of the Gas Bureau of the city of Philadelphia, where the loss from leakage in the loss of amazonatelphia



Fig. 82 Exterior of a pape impared to obstacl are Springfuld. III

for gas, for a period of ten years, some \$5.750 ones, and for the years, ISOI to ISO5 averaged two millions of order for years day.

The Brooklyn Union Gas Company (Brookly) & A has about 700 miles of gas-mains of all object and 28th rade of we wield motive as service-pipes. The latter and their latteries are forced to be leadly corrected wherever uncovered for examination. Hertroleus from stray electrical currents is manifest in manifest in a dec. Thirts eight service pipes in one street block were completely destroyed in three years. The east-iron mains are reported to be convailly an a good condition so far as electrolytic action is conserved to succeed to the converted as a few cases in what is called "the damperous objects of that is, in the vicinity of the electric station power hospect.

The loss of gas from all of the antierground of tenso as there etc. in 1899 amounted to 13 per cent of the total or religiously 1 and

corroded to such an extent that the whole pipe-line had deteriorated 50 per cent in four years. The voltage in this line was from 2 to 9 positive. At a voltage averaging 4.5, a 6-inch pipe became useless in five years.

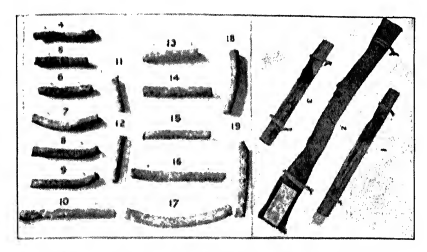


Fig. 83 - Lead water service pipes and telephone calde coverings in Brooklyn, N. Y.

Water mains that were laid and tested to withstand a pressure of over 300 pounds to the square inch, at the end of four years leaked at almost every joint at 150 pounds' pressure. The voltage was 4.5, and the lead joints were badly corroded.

Electrolysis of water paper at Kansas City, Mo.,* Mr. G. B. Wing, Superintendent of the Metropolitan Water Works, reports that specimens of the scal at a distance of two inches from some of the corroded pipe showed 4.67 per cent of iron, and at a distance of one foot, 2.65 per cent.

The corresion of wrought iron and lead pipes was more rapid than that of east iron pipes; the amount of corresion in all cases depended upon the impressed of the current.

and to the elevated railway structure, there was a difference of from 2 to 2.5 volts. At one-fourth of a mile away the current all passed to the elevated railway structure, ran a fourth of a mile, then returned to the water pipes and changed again to the railway structure in about half a mile.

The same conditions were found to prevail on an opposite section of the elevated railway structure, extending for about a mile in the opposite direction from the trolley terminal.

The difference in potential between the elevated-railway columns

and the street trolley rails and water-mains ranged from 1.30 to 1.50 of a volt, and indicated that the current came from an electric lighting station. It was also shown by the tests that a trolley-line using the rails, water and gassimins for its return service, can spread the corrosive influences for a mile in either direction through the various subway conduits, pipes, and elevated-railway structures; also that the conductivity of a 50 pound street or tee-rail is about equal to a copper real 1 inch in diameter, or tive No. 000 B, and S, copper wires.

Tests applied to the Brooklyn Bridge suspension cables showed

Tests applied to the Brooklyn Bridge suspension cables showed that generally there were 3 volts positive to the rails of the trolley railway on the structure. The effect of these currents upon the anchorages of the bridge led to a number of tests of the upper ends of the nucleorage metal. The tests are believed by the bridge engineers to show "that no damage such as might be expected from corrosion of underground metal has thus far taken place."

A wise distinction between corresion and electrolytic action. Had the question of the corresion of the rails in the street tracks been put to the tridley railway engineer corps, they would probably have been positive that no such corresion was present or possible; in fact, they were indifferent to or ignorant of the corresion until the examination by Mr. Ixandson.

It required the public evidence of a half-dozen of broken suspension rods and panels of sunken radway tracks to convince the Brooklyn Bridge engineers that a serious case of neglect and corrosion exists in the structure, and had progressed far enough to be dan-

Iron and steel bodies exposed to conditions similar to bridge anchorage metal have been found badly corroded within a few years after being placed in position, and there is no reason to infer that any bridge anchorage will be an exception.

The rate of corrosion from natural causes has been fairly determined. In anchorage work this will be increased by any electrical currents that may reach them, and it is inevitable that they do reach them, and no means of preventing it now exists.

The decay of metal by electrolysis has been approximately ascertained. The escape of the voltages and amperes used in street railway service is twenty times that necessary to induce ferric corrosion and often more than twice as much as is necessary to decompose water in mass.

A current of 0.3 ampere is sufficient to corrode a lead covering to a cable or the lead in a pipe-joint. Electrical engineers report cases where the lead covering of cables has been destroyed in six weeks after laying.

A potential of ${}_{16^{10}6}$ of a volt is all that is required to induce ferric corrosion two miles from the dynamos.

A difference in voltage of 20 volts has been found between the two ends of the Brooklyn Bridge cables, and the difference in voltage ranges from 0.75 to 3 volts at all hours and at all times in the day whenever tested, and is always found electro-positive to the ground.

So long as electricity obeys the known laws pertaining to its generation and transmission, it will select the line of least resistance, though it may not be the shortest in returning a major part of the current to its individual source of generation. It will also divide en route, pick up other electric currents in the most erratic manner, and deposit them in unexpected places, generally inaccessible for observation or repairs.

The large amounts of voltage and uniquees used in railway-motor systems render stray electric currents more certain and electrodysis more constant, even if a "shunt" of the current from any adjoining bridge cable or structure were possible. At the present state of the

times in the earth and air and never, or but momentarily, of the same potential.

Hundreds of electric installations of a great diversity of power surround these bridges and provide a cause of danger that at present, if known or suspected, has no remedy or safeguard.

The future results of electrolysis on all suspension bridges may as well be recognized now, rather than be left till the inevitable catastrophe befalls,

The corrosion of ferric bodies, not aided by electrolysis, is known to be progressive, being nearly 50 per cent more the second year than the first, and so on for each succeeding year.

During the construction of the Britannia Bridge over the Menia Straits, some rejected plates 176 and 7 inch thick, were left unprotected and exposed to the spray and wash of the sea. In two years they had corroded so that they could be swept away with a broom.

A few pieces of ironwork embedded in mortar or walled in some ancient building, or an old water-gate here and there, in some very favorable situation, may have remained uncorroded, but there is little magnestionable proof that iron or steel in the form adopted for bridges or structural frame-work will last more than two hundred years.

In the Niagara Falls and the Alleghany River suspension bridges, after about twenty five years of duty, an inspection showed that some of the wires in the outer strands of the cables outside of the anchorages were corroded through, but the second and interior wires were sound. The reason assigned for the corrosion of the outer strand wires was that the "creep" of the individual wires under the varying strains due to the load and constant changes in temperature had worn away the bailed lineard-oil and other coatings applied when the wires were strung and allowed atmospheric moisture to reach them.

It is now proposed to abandon the boiled oil or paint coatings of the cables and to use a mixture of vaseline and plumbago. When the wires are strong and ready to bunch into strands and cables, all of the interstices are to be filled as far as possible with this stiff undrying mixture, that is to act as a lubricant for the inevitable creep element of weakness in providing a foundation for the paint contings that are supposed to seal the cables water tight.

Mr. Robert Mallet, C.F. (Dublin), made a report to the British Association in 1858 on paints for bridge and cable iron work; "That he had tested ten of the best and most reliable term points and varnishes then known, and not one of them remained adherent and undecomposed for a single year under water. In most air, and under conditions resembling English seasonst fog, their state was not much better. The presence of mosture even to the extent of a partial saturation of the air, developed a fungus, the decomposition of which was almost as fatal to the life of a paint as immersion in sea-water."

Government authorities state that there are over 300 suspension bridges in Europe, of a great variety of spans and industrial importance; many of them having evelors instead of wire cable suspensions. It is also stated that the life of the wire cables and anchorages have been found to be precarrous, for oxidation was in progress in the interior of the cables, while the anchorages were weakened from the attack of some element "not at present defined" (evidently electrolysis). That there was no reliance to be placed on the preservation methods, or any certainty that they had effect on the life of the structure beyond twenty-fixe years.

The failure of the Anglers were cable suspension bridge showed that it was impossible to keep the hydrate-of-line coating used there in immediate contact with the anchorage metal. Moisture and earth neids, carbonic acid from the atmosphere reached the metal, and the line-coating was practically useless to prevent corresion.

M. Bernadeau in the "Annales des Pents et Chausseur," 1881, refers to a bridge in which of the 150 wires forming a cable only 15 were in good condition, the rest were brittle as glass. The bridge had been in use less than forty years.

Two other suspension bridges of short span fell after twenty-six and twenty-eight years' duty,

The suspension bridge over the Ostrawitza River at Mahrisch-Ostra, finished in 1851, failed in 1886 from a fracture of one of the The anchor-bars in this bridge consisted of 12 links, one of which was completely corroded away and the others were reduced to about one-sixth of the original size. The original sectional area of the anchors was 24.4 square inches, but had corroded to about 4 inches. An official examination and report of the strength and condition of the bridge was unde in 1885, one year before its failure, which stated that "The bridge has been examined in all its parts and is in good and safe condition." A squadron of Uhlans went down with the bridge.

An exmoination of the wire cables of a suspension bridge, where coal-tar and line had been used to coat the wires, also to fill the interstices between them where the cables entered the anchorages, showed that these cables were wrapped with \(\frac{1}{0} \)-inch diameter wire and then a canvas jacket saturated with coal-tar and line placed over them. After less than twenty years' duty the tar had partially decomposed and disappeared and the cavities were filled with a dirty, grayish liquid. The wrapping wire, also the seizing wire on the strands, were in many cases rusted through, and the cable wires deeply pitted. The damage to all the wires extended about three feet upward and outward from the cable anchorages. Beyond this there was a little rust, but no pitting, and still further from the anchorage the paint on the interior of the cable was gammy and undried.

*French engineers of reputation now prohibit the use of white-

lead or any quick-drying paints on anchorage cables. The failure of a number of suspension-bridge cables in France was directly traceable to the use of that kind of paint. Chalking and cracking of the coating, owing to the ceaseless changes of temperature to which they and the cables were expected, admitted water and held it, and corrosion of the wires at or near their lowest position in the cables was the result.

Emphorbium paints possessing elasticity, tenacity, and a quality that prevents them from drying bone-hard and becoming brittle, have proven the best paints used by French engineers for cable or other ferric work. Emphorbium being of a non-corrosive and antifouling nature, prevents the growth of atmospheric fungus, the decomposition of which produces an acid highly corrosive to iron.

Other resingue or varnish paints dried hard and brittle and soon

to corrosion or electrolysis and not to overloading, shows the imperative necessity of having the cable and particularly the anchorage metal accessible for inspection at any time. Controlled of cables and metals in the air may be in part observed, but electrolysis occurring in the lower and hidden parts of the anchorage, and not reparable nor preventable, should be guarded against by a metallic connection at the anchor-plate end, that will lead off all electric currents and extinguish them in the earth and not in the metal of the structure. This connection can be renewed when corroded, and all shourts or attempted cut-offs of the current above the cronnel line avoided. In the case of divided currents, cut offs have been bound to be unreliable.

Coment coatings or concrete cannot retard electrolysis if any moisture is present. Mr. Eiffel found the iron tag bolts placed in fortification musoury two lumides! years are, had enlarged from two to two and a half times their original diameter by ricting in the mortar in a dry location.

Rust once established, carries within itself the elements for a conscless life. Even in a place battle rust begets in to Hydrated rust carries over 20 per cent of moisture, and so long as it can attack a fresh surface of iron and east off the thin files of excite as it forms, it will release enough oxygen to begin another excless action

Farraday's law of the correspon of metal in weak accludated solutions and electrical energy applied to the anode i. I titl's etains of iron per square foot, per ampere hour. The late of any ferric body can thus be approximately ascertained before its consistence. The methods and means for its preservation should inverse the rice tearerful consideration of the engineer and other responsible for its preservation. That its preservation is more shiftened than its planning or construction does not remove this serious responsibility.

The borough of Brooklyn (New York Little Law does to the collection)

The borough of Brooklyn (New York Utty) has about 800 miles of water-mains of all diameters from 1 to 18 males. Many of these pipes in the early construction of the water-works were east in takegow from a firm close-grained east iron which showed a white surface on fracture, indicating a large amount of construct cathon in the metal. Many miles of these pipes were also contest with the

were hid, also to the composition of the cast iron.

Scotch gray or forge iron was thought to be exempt from corrosion, and the Roebling bridge anchor-plates were made from this brand of east iron and placed in the anchorages under this idea. But many of the pipes were east in American foundries from American east iron, and but little difference in their corrosion and that of the Glasgow-made pipes had ever been noticed. After the effect of corrosion by electrolysis had been noticed, to determine whether the composition of the east iron had any power to prevent it, pieces were ent from the foreign and American east-iron pipes, also from soft east iron containing but little combined carbon and more graphite than the Scotch irons, and used as anodes in various electrolytic cells. The electrolytes consisted of samples of earth from various parts of the city, moistened with distilled, hydrant, and sea-water. The cells were exposed to the action of currents of different voltage and amperage.

In every case the anode was corroled, showing conclusively that there is no immunity from electrolysis of east iron used for water-pipes, because of its chemical composition.

It was determined by the observation of the water-works' engineers that the tubercular corrosion on the water-pipes when unprotected other than by the usual thin coal-tar or bitumen pipe dips was at the rate of about $\Gamma_{0,0,0}^{A_{0,0}}$ of an inch yearly, there being a difference in the rate of corrosion in the pipes of different metals, markedly in favor of the close-grained, white firm irons.

Summarizing the report of many other water-works' engineers, it appears that corresion from electrolysis, tubercules, or from other causes is more rapid in wrought-iron than in east-iron pipes, irrespective of the kind of soils they are buried in or to whatever influences they may be expand.

The small amount of electrolysis in the city of Brooklyn gasand water mains was finally attributed by Prof. Samuel Sheldon* to the presence of the hard, thin, vitrous scale formed on them at the moment of easting in green sand molds, and that this scale was a non-conductor of electricity. This scale is similar to that noted on to on page 328.

A piece of the sand-conted pipe was covered with an insulating paint, but leaving exposed a small area of the alwate conting. This pipe was made the anode in an electrolytic solution. Lass current flowed through the solution under a given ICM 1 - than under a cular conditions with an anode from the same piece of payer, but expecing a clean iron surface of the same area to the same entrents. In some of the experiments, no current at all parced the scale until the voltage was raised to a number of older. The water pures conted with Dr. Angus Smith's compound appeared to be less affected by electrolysis than the pipes not readed. The moulating quality of the compound added to the power of the silicate conting to resist the stray electrical currents of law pedential. In certain distincts of the city where high potential currents reached the paper scheetrolysis was present, but concealed by the time and unlocken reating of Dr. Smith's and other heavy anti-corrective coating and was the more dangerous on this account

In every city in which electric street radway lighted, and power service is developed, there will be a number of district some pending to the number of power stations and plants for in isomed electrical generation. Each one of these installations will struct currents from its own district that can generally be very slovely defined in the ordinary working of the station. But these houselative become very irregular, daily and hourly, from the varying nature of the currents required for the work to be done in them tespertisely. A difference of potential has been noted as high as 40 volts between sufferent points in the same district or between adjuming district. Hence these boundaries are always shifting to a preater or be existent at all times. It matters but little from which district the a irrest teacher underground metal or what its potential, electrols reachers a correspondent metal or what its potential, electrols reachers are every case.

Where electrical installations of a known amperance of litters to 50,000 are in daily use, there will meastable be once a leakage of the direct and return currents as well as a contain energy to the archiection currents, always present for the correspond of a conal substitute to currents.

alarmst, the plant tacts may as well be recognized now as hereafter, when the particularly dangerous character of stray electrical currents of low voltage and large amperage is forcibly presented to the public in the sudden collapse of some important structure or gas-and water supply systems

It is a false reliance that masonry, mortar, concrete, or cement are impervious to moisture and incapable of acting as an electrolyte such as would induce electrolysis. They are not insulating substances, or at the best only in the smallest degree under the most favorable circumstances. They are positively porous and in nearly every case, whether tested in large or small mass, are permeable to all waters or moisture and gases, and in but a few exceptional cases ever become thoroughly dry.

*A number of electric light cable conduits in Paris were constructed.

of concrete, particular care being exercised in the selection of the hydraulic cement and sand used, as well as the ramming of it into place. The conclust were far above the water-line of the city's soil and were considered to be water-tight. The copper wires soon became covered with verdigits and copper chloride and so reduced in area that grounding and heating were of frequent occurrence from the normal currents of the service. A number of minor explosions also occurred, due to the gases formed by the decomposition of salt water that filtered through the cement when salt was strewn on the roadway over the cable condint to melt the snow. The gaseous mixture contained oxygen, hydrogen, and chlorine, the latter gas being due to the chloride of sodarm in the salt water. The leakage of the current furnished the electrical energy to decompose the salt water and that was deposited upon the copper wires.

Farthenware conducts were also used, but were not water-tight, and the same decomposition of the salt water and corrosion of the copper wires occurred as in the concrete construction, and their use was soon abunchered. The electric cable wires are now taped or covered with a bituminous compound to prevent electrolytic action.

President Learnest, in his innugural address to the New England Association of the Engineers, March meeting, 1902,† stated that the conclusions derived from a large number of to be and observations of the effect of electrolysis on the gas paper vertical in a number of cities in New England were: "That gas paper land with lead joints have 15 per cent greater resistance to electric current, than waterpipes of the same diameter with similar points. Screwed joints in wrought-iron pipes have about the same resistance as the lead joints in cast-iron pipes of the same diameter. The resistance of a Port land-cement joint, as ordinarily made, was from 15,000 to 20,000 times the resistance of a lead caulked joint comparing paper of equal diameter. That the resistance of the cement joint depended in a great measure upon the amount of moisture that the cement takes up after setting.

"The conclusions drawn from the experiments tande on gas pipes of all diameters laid in short or long sections were. That all possible resistance should be inserted in the pipe mains by making the joints of some insulating or semi-insulating material, with an arbestos or tar-paper ring between the abutting ends of the pipe in order to break up the pipe-line into as many metallic units as possible, and isolate them from all other pipe or trolles systems, so far as practicable and mechanical conditions would allow. The pipes should be control with a water-proof compound. The ordinary toundry dip conting or painting with oil paints had no appreciable effect to delay or diminish electrolysis of the pipes. Neat his laudic centert to delay or diminish electrolysis of the pipes. Neat his laudic centert coverings were worthless on account of the papers actual of the pipes by frost or mechanical injury.

"A covering made from 3 parts of dry clean sand and 2 parts of coal-tar boiled to a pitch at 660° b, made a maxture that was shightly clustic at ordinary temperatures, was thoroughly water proof and when applied to the pipes to a thickness of 13 melocs, had an inculating resistance of over 1 million of ohms to the cube melo. A short piece of 2-inch pipe covered with this maxture and immersest in a strong solution of water and soda ach for 8 hours, showed no signs of the absorption of any of the solution, nor had any electrone of

The corresion by atmospheric exposure in the Brooklyn Suspension Bridge superstructure is deeply scated in every square foot of the structure and is beyond correction except by rebuilding it. It takes a corps of painters con tantly at work three years to paint the structure, and the reating principally serves to mask the corrosion. The voltage of the electrical currents passing through the suspension cables has been reterred to, but their corrosive effects upon the anchorage metal from the macrosolubry of the lower ends must always be a conjecture. The e-patts are beyond repair, and the electrical currents, whether from the trolley railway on the bridge, or from the scores of large installations surrounding the structure, are of large amperage, constant and uncontrollable in action. That they will not prove active agents of electrolysis is not in accordance with

In the other hast River suspension bridges, where steel instead of nonsours parts are used to carry the suspension cables and the superstructure, it is expected that the large metallic contact of the wire cables at the top of the steel towers will form a short circuit and ground for any electrical energy that may reach them from the railway, instead of using the anchorage metal for a terminal. This theory can only be determined after the bridge railway has been put in operation. The many points of praction of the bridge trestles with the ingreation. The many points of praction of the bridge trestles with the ingreation parts on both sides of the river may divide the trolley currents into a number of short circuits so as, in a measure, to protect the structure. But this cannot prevent the currents that come to the anchorage metal from the installations surrounding them, from using the valides for their transmission, as they provide the last and shortest metallic path for their ceaseless circuit.

A further fast relative to the electrolysis of the uncharge metallies in its condition even before any material strain other than the weight of the Tost path calder cause to it. In the Williamsburg Bridge, the anchorage puts were carried down into the solid rock and near if not below the water level of the river. These were not scaled water-tight by carry effectual method in laying the superincumbent massure. The anchorage metal was put in place under a continual with Smith & Co.'s "Durable Coating," the advisability of applying a baked japan coating being disproved on account of its cost.

When the chain of cyclars were ready for the cable construction, an analogous sit was parameted dry and the cyclars increased. The coat

an anchorage pit was pumped dry and the cycliars inspected. Though the bars were covered with two coats of "Turable Coating," and in place only about two years, the paint was nearly destroyed, and corresion over the whole surface of the bars wherever the water had reached them was virulent. The broited mean in the pit and the close association of the cycliars together and to their bad, rendered the cleaning and repainting of their difficult and in many feet of their length impossible. A new cost of varnish paint, however, was applied in the damp atmosphere of the pit.

For the future protection of the exchara it is projected that when the bridge is completed and the chain of bars are hearing their load and have adjusted themselves to their permanent position, to paint them again and fill in between and around them for a foot or more with melted bitumen and to fill the pit with concrete. This plan does not reach the anchorage plates and metal in the lower end and mac cessible part of the pit. The corresion scripent is only scotched (not killed) and will remain imactive for but a short time, or only so long as the concrete remains thoroughly dry, something impossible to maintain. The bitumen conting insulates the exchara so far as it can be

applied and passes on whatever electrical currents may reach them from any source to the lower end of the anchorage, where the inetal is not protected, corresion in progress, and imposition almost if not quite impossible. To think that electrodesis will not take place in the metal at both ends of the bridge is to agree facts already established. Electrodysis, or even corresion from the contact of metal with moisture, in this case, is not the sin of the paint manufacture; where to place the blume is not hard to find

The introduction into marine service of appliances for the generation and use of electric power and light has slewelequed a new field for electrolysis, that scrionaly endangers the efficiency and life of all vessels so equipped. An examination of the United States exames Breakly a for the purpose of electronicing the effects of a recent grounding of the vessel, has recorded the fact that the test of a recent grounding of the Electrolysis on board a steel ship is not unlike the same development by direct or stray electric currents in land or underground structures. Wherever a current of any potential leaves the metal, electrolysis is the result. In the case of naval vessels there is an enormous amperage present at all times and that cannot be returned to the dynamos, even with an increased capacity of the return-current wires over those employed for distribution of the current.

Divided and induced currents, also the electric energy developed by corrosion itself, will seek their own course either in returning to the dynamo or extinguishment in the ground connection. The latter, in the case of marine work, being water saline or foul in character, is a more efficient electrolyte than any earthy substance. Hence electrolysis in marine constructions will naturally be more rapid and virulent than on a similar ferric area and current exposure on land or in underground structures.

As before stated, no paint or plastic coatings of the metal will prevent electrolysis. At best they may temporarily mask its progress, but it only requires a short time or a slight change in the conditions to reveal it.

Corrosion or electrolysis of marine metal can only be controlled by the use of some alloy of steel that will minimize the action, and by such an increase in the thickness of the parts of the ship exposed to corrosive influences as will for a time provide for any reduction of strength in the corroded parts or alloyed metal; also by a plan of construction that recognizes the possibility of the evil and provides that corroded members can be removed without practically rebuilding the ship. (See page 339 for Dr. Wurtz's protective method.)

Insulation of motors and their connections and the positive prohibition of the connection of any electric current, even of the smallest amount, to any part of the structure will reduce but not prevent the dangers of electrolysis, whether in marine or land constructions.

In either case, constant and thorough inspection of all ferric surfaces by an inspector who knows what to look for, and is competent to recognize it when it is found, is an essential. Even if the inspector cannot avert corrosion when found, at least the danger can be noted, watched, and a warning given when it is time to desert the ship. All of these essentials appear to have been absent in the